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**Cheng et al.**

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(54) **TRANSGENIC FISH AND USES THEREOF**

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(21) Appl. No.: **13/769,533**

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(22) Filed: **Feb. 18, 2013**

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(65) **Prior Publication Data**

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**Related U.S. Application Data**

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(62) Division of application No. 12/730,956, filed on Mar. 24, 2010, now Pat. No. 8,395,018.

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(51) **Int. Cl.**

**C07H 21/02** (2006.01)

**A01K 15/00** (2006.01)

**A01K 67/027** (2006.01)

**C12N 15/63** (2006.01)

**C12N 15/85** (2006.01)

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(52) **U.S. Cl.**

CPC ..... **A01K 67/0275** (2013.01); **C12N 15/63** (2013.01); **A01K 2217/05** (2013.01); **A01K 2217/052** (2013.01); **A01K 2217/203** (2013.01); **A01K 2217/206** (2013.01); **A01K 2227/40** (2013.01); **A01K 2267/0393** (2013.01); **C12N 15/8509** (2013.01); **C12N 2800/106** (2013.01); **C12N 2830/002** (2013.01); **C12N 2830/008** (2013.01)

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(58) **Field of Classification Search**

CPC ..... C12N 15/00; C12N 2830/00; C12N 15/63  
See application file for complete search history.

(57) **ABSTRACT**

The present invention relates to a transgenic fish having at least one genomically integrated expression cassette containing a 5'-regulatory nucleotide sequence responsive to hormones, particularly estrogenic hormones, connected in a functional manner upstream of a nucleotide sequence encoding a reporter protein. The present invention further relates to methods of using the transgenic fish for various purposes, including, for example: (1) identifying estrogenic endocrine disruptors; (2) monitoring estrogen-like activity of test samples; (3) identifying anti-estrogenic endocrine disruptors; and (4) investigating the effects of endocrine disruptors on liver regeneration. Expression cassettes, host cells, and transgenic cells of aquatic animals are also disclosed.

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**20 Claims, 15 Drawing Sheets**

Fig.1

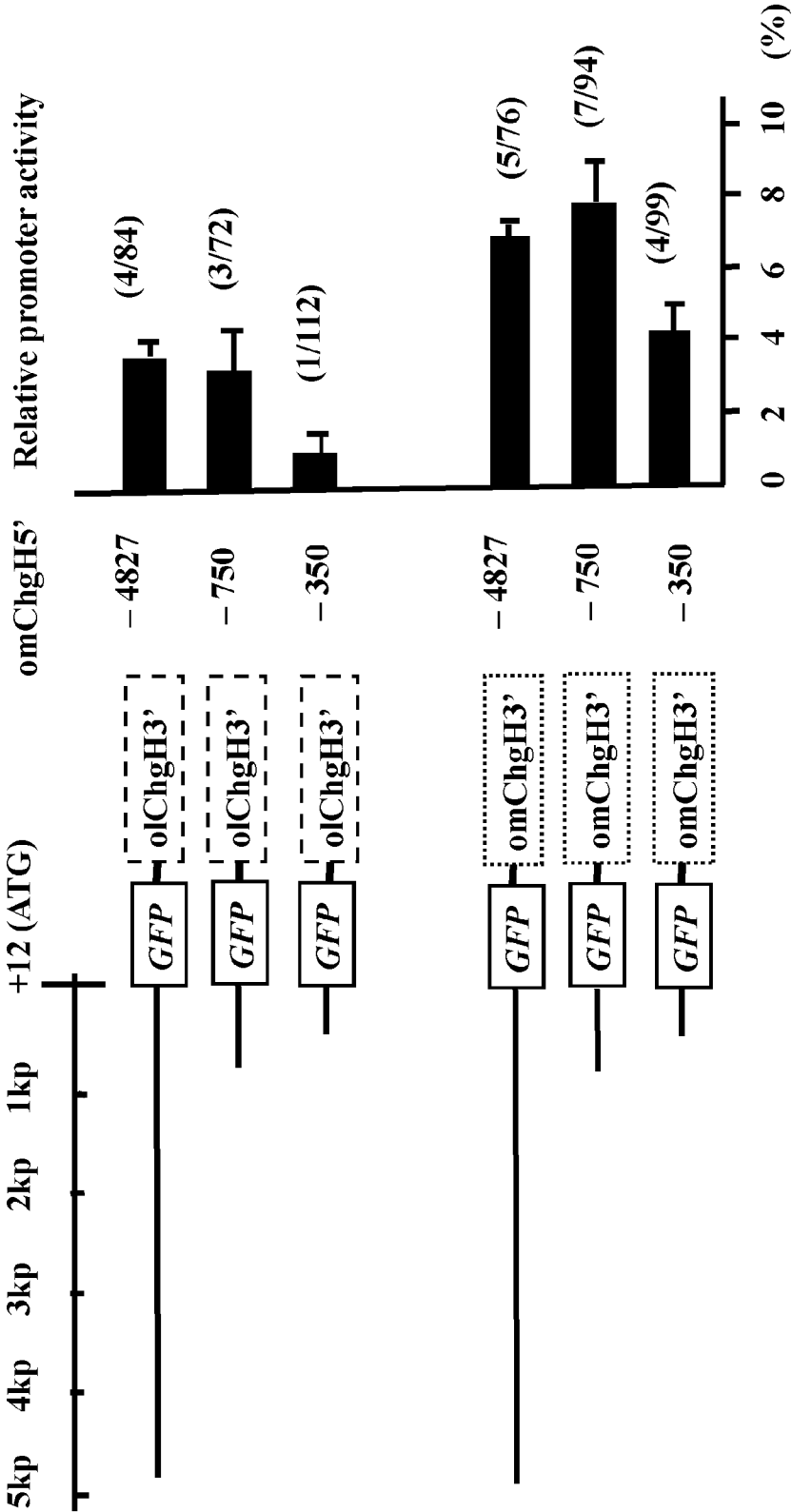
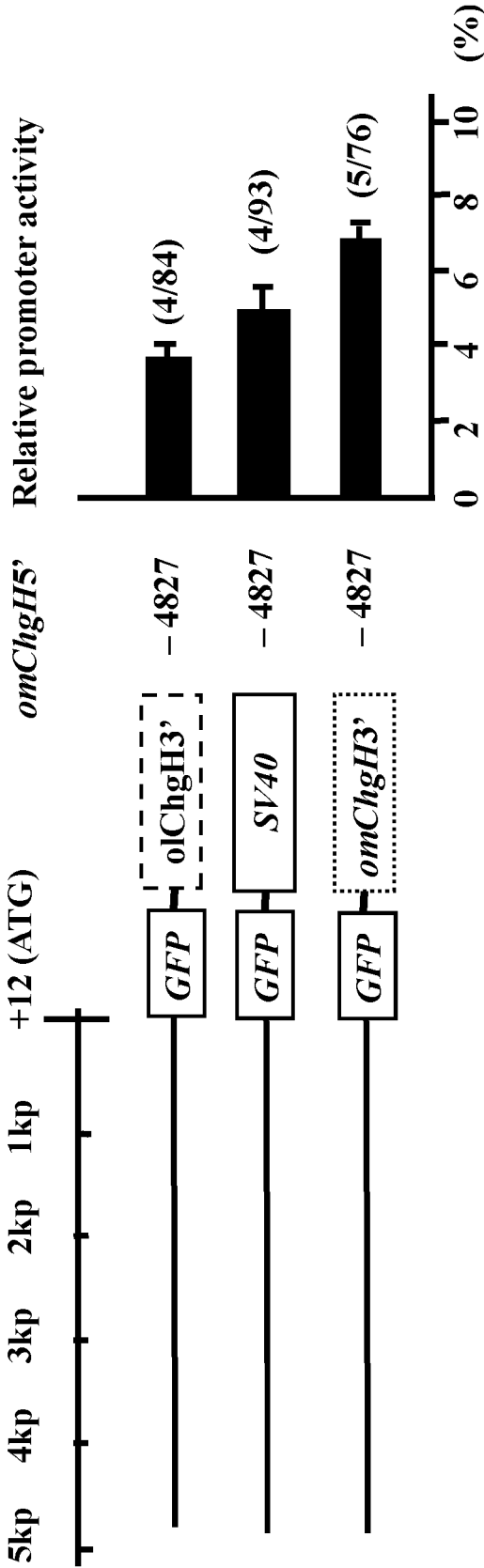


Fig.2



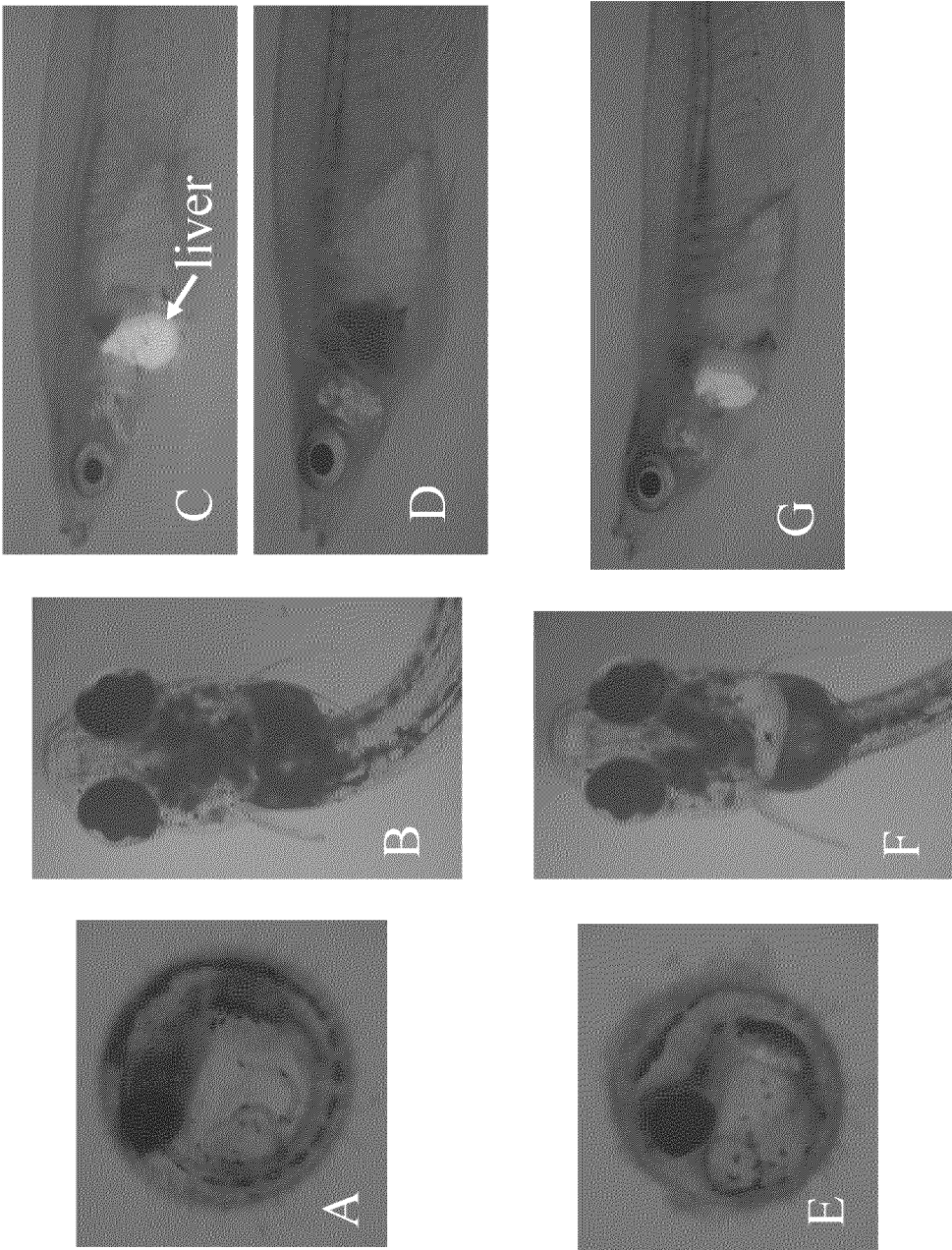


Fig.3

**Fig.4**

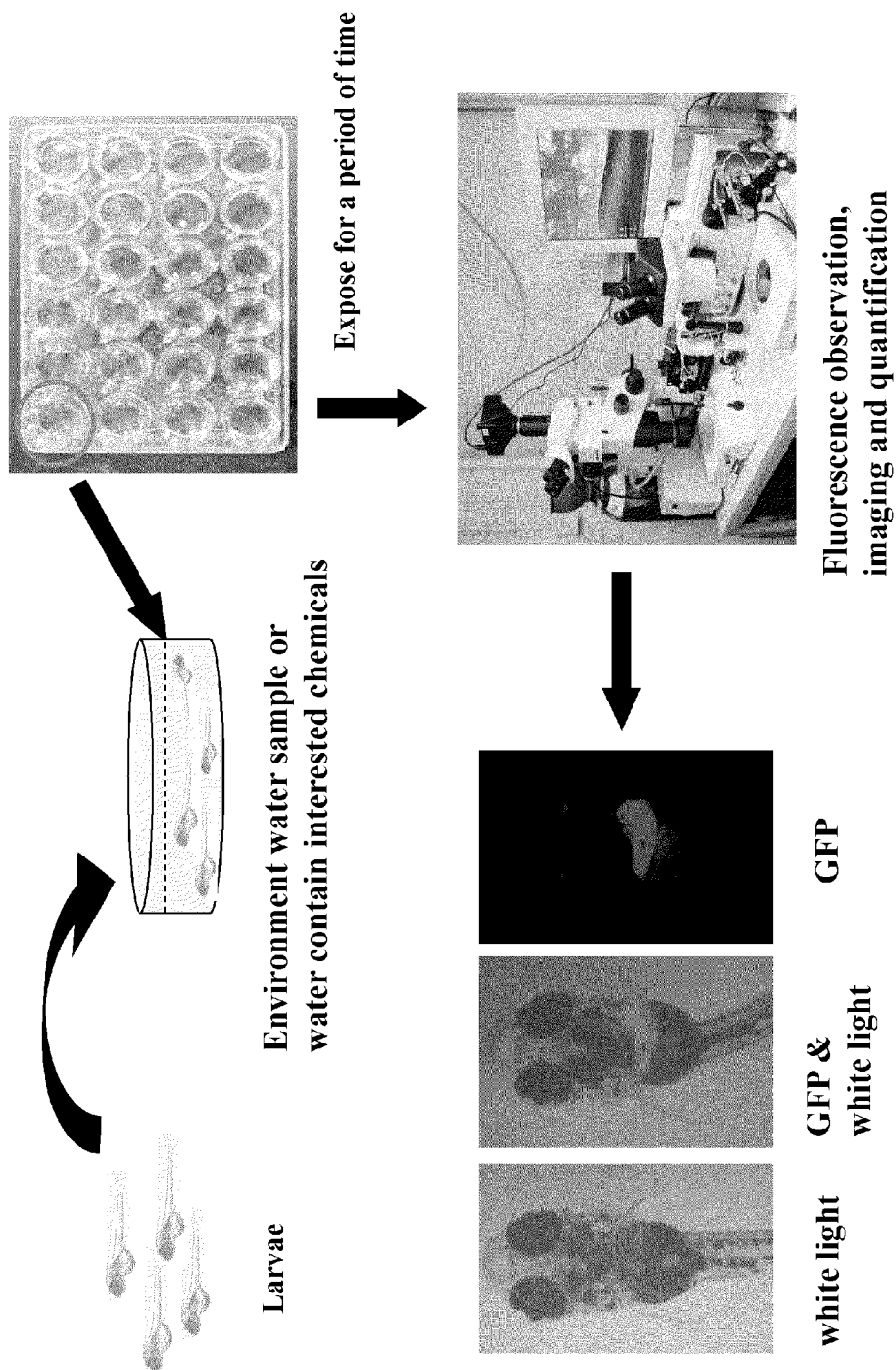
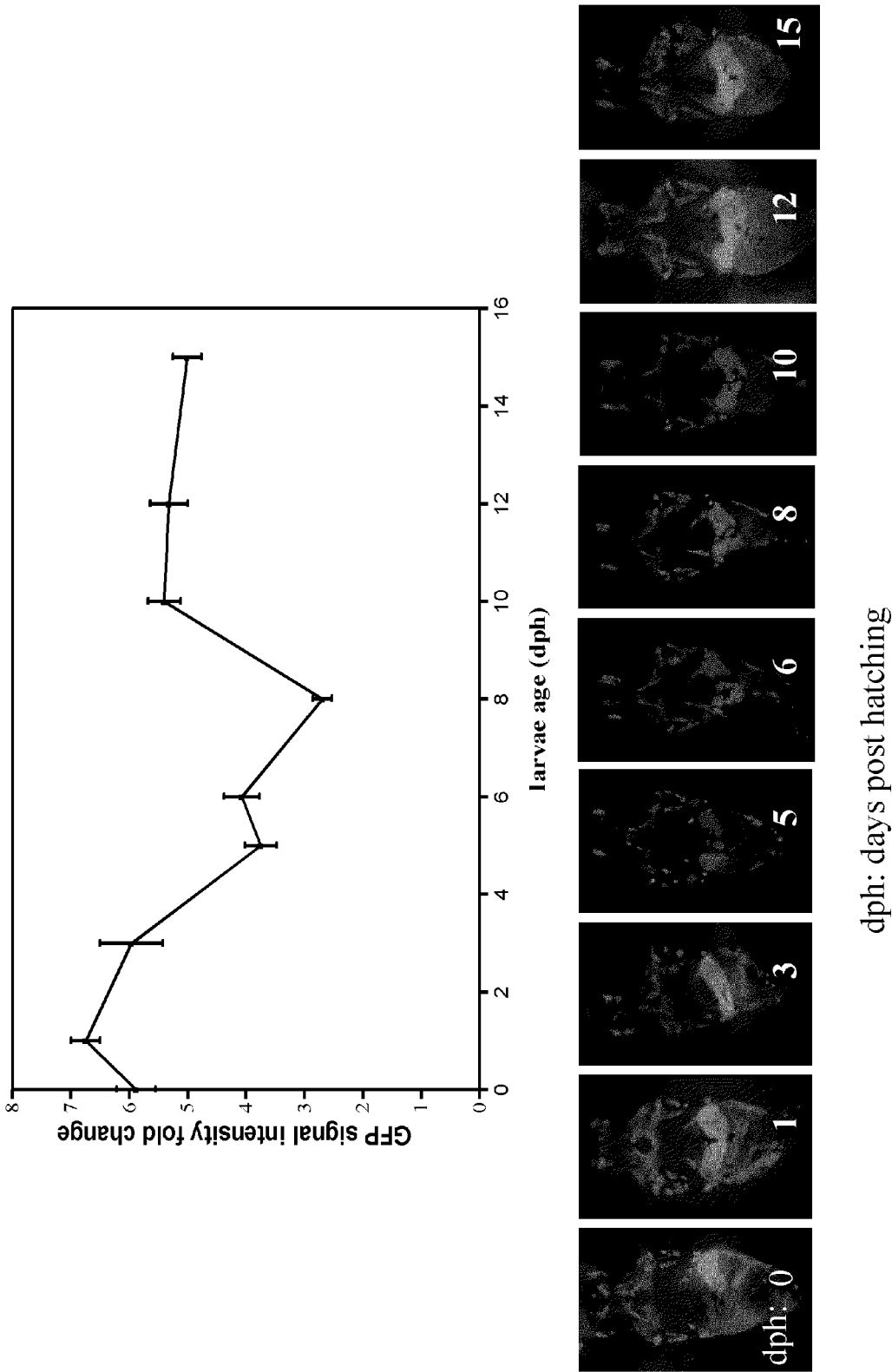


Fig.5



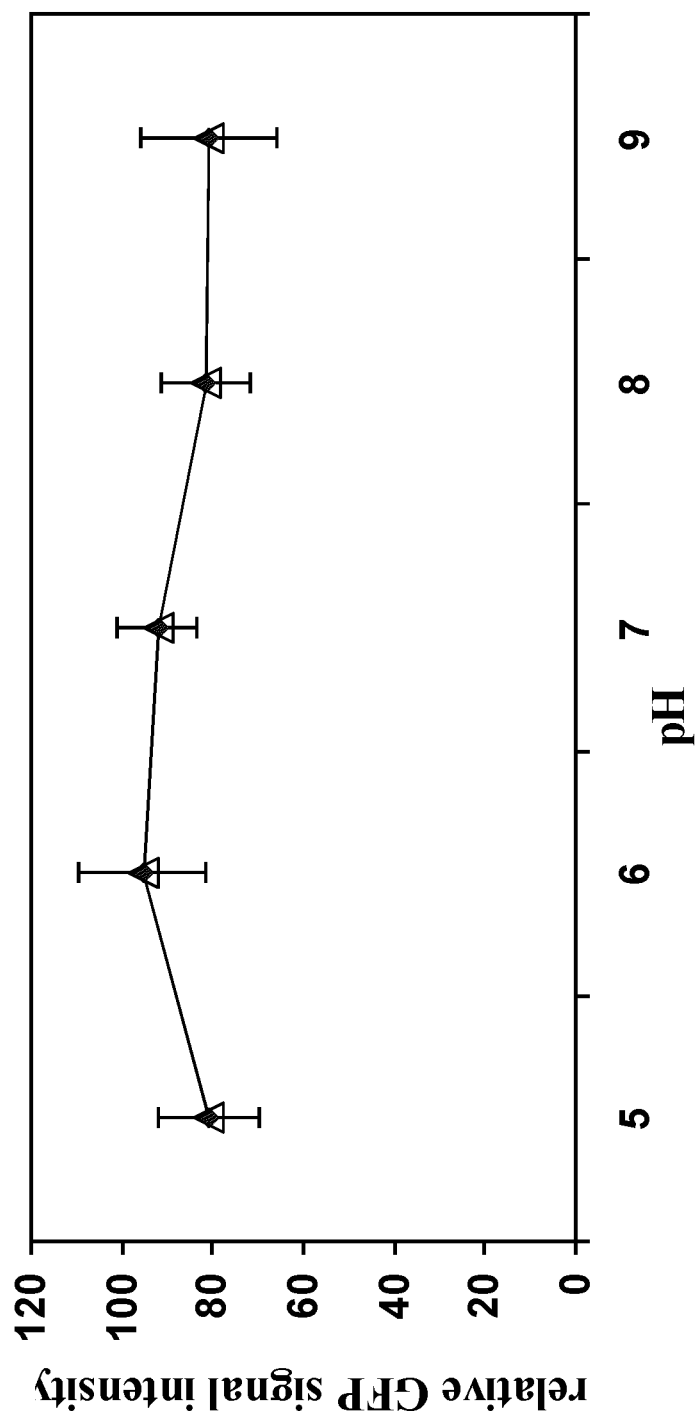


Fig.6

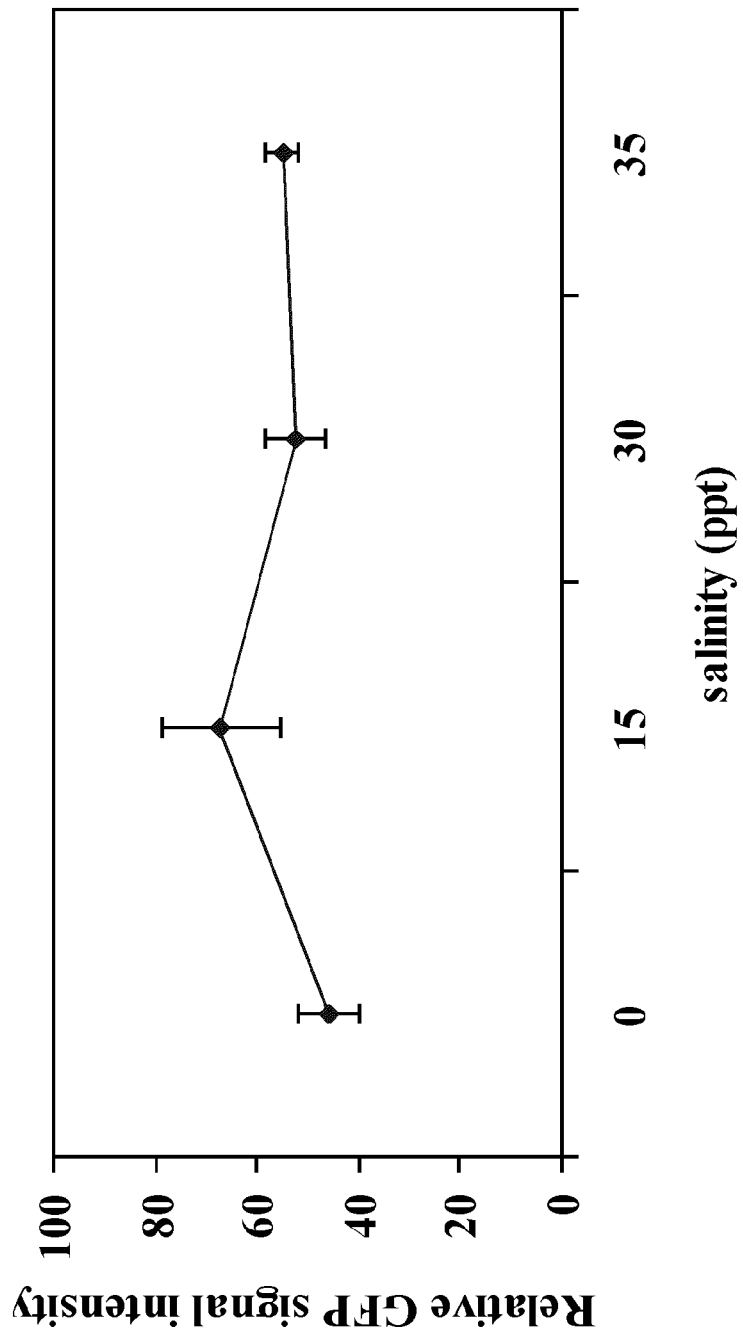
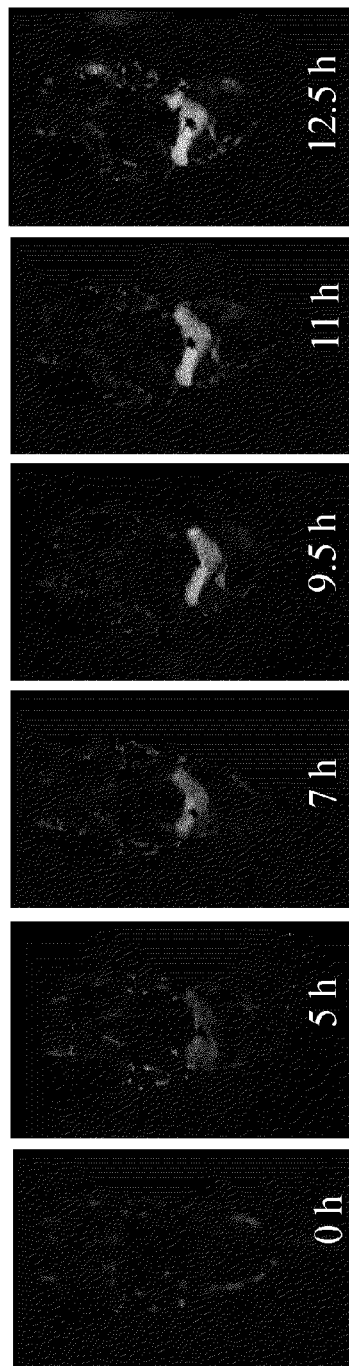
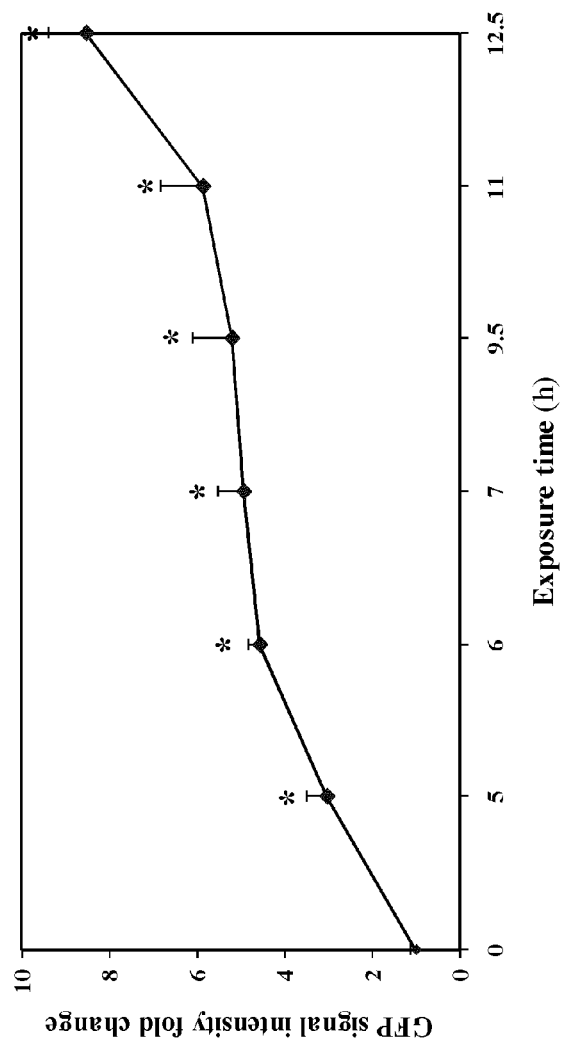
**Fig.7**

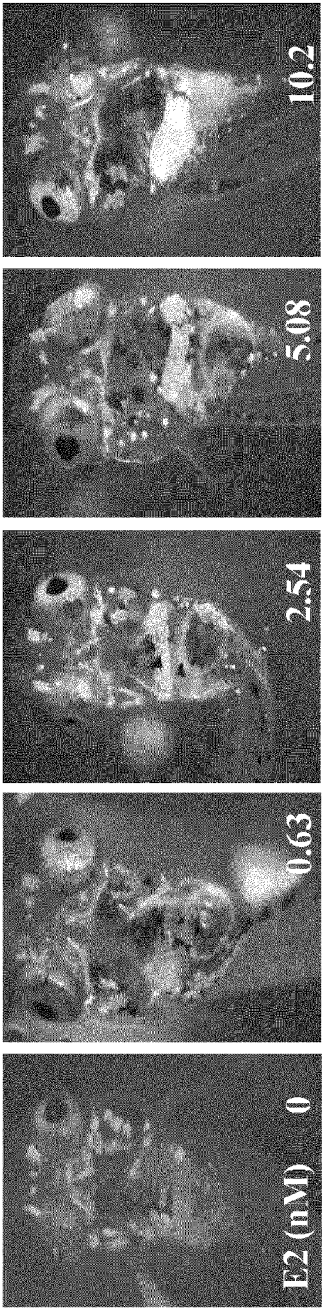
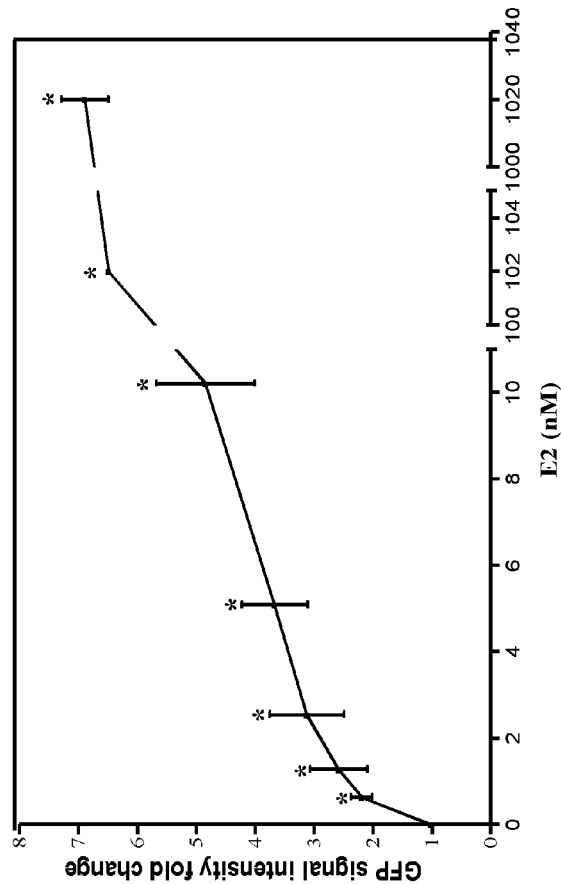


Fig.8



\* $p < 0.05$

Fig.9



\* $p < 0.05$

Fig.10

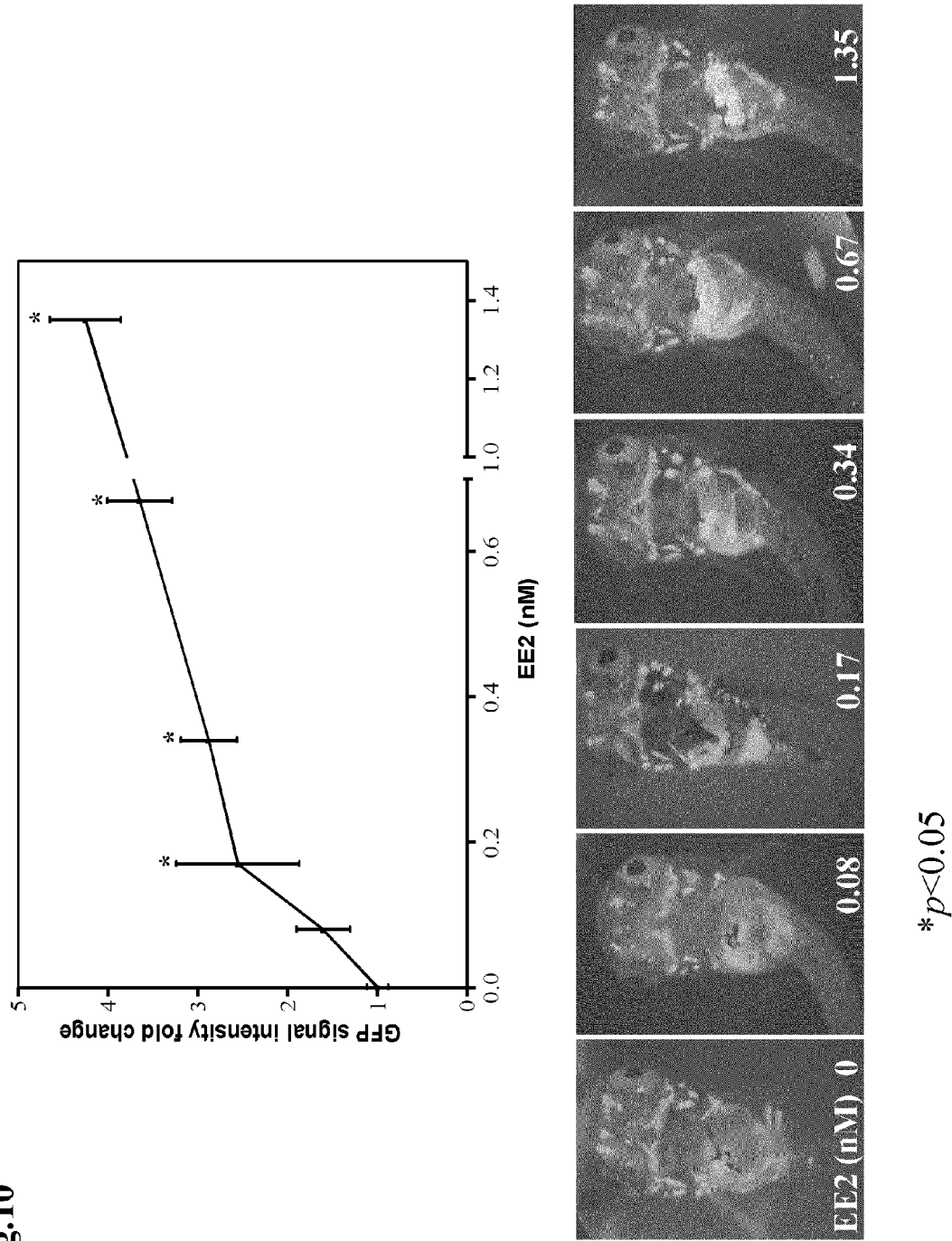
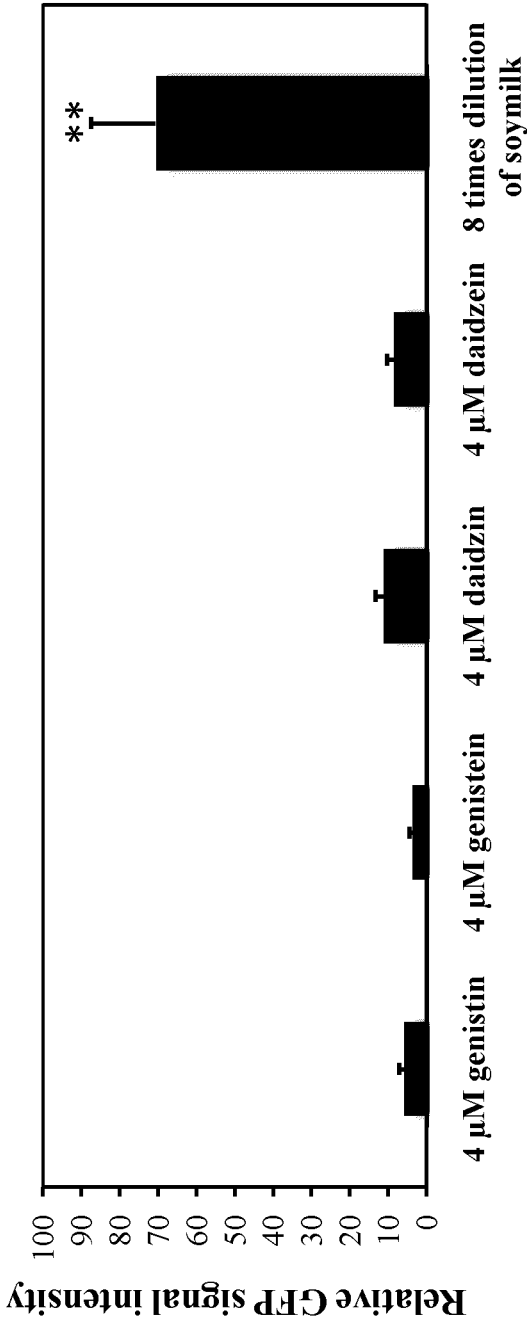


Fig.11



**\*\* $p < 0.005$**

Fig.12

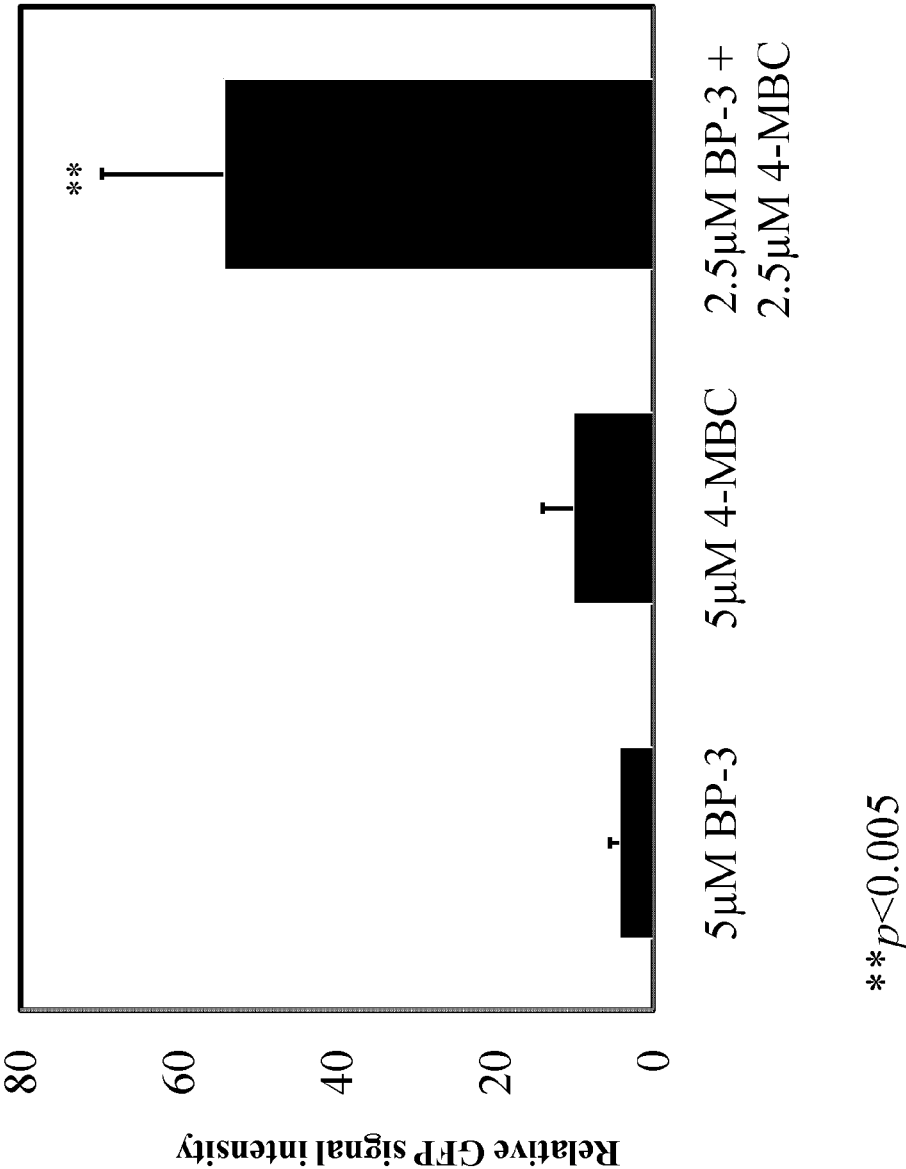
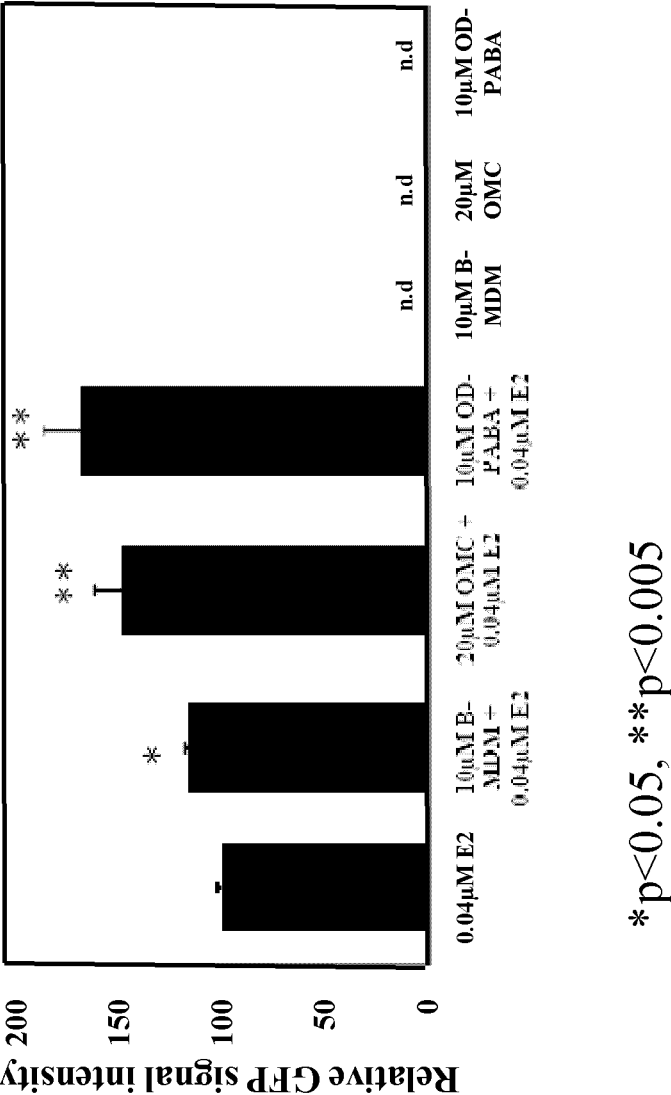
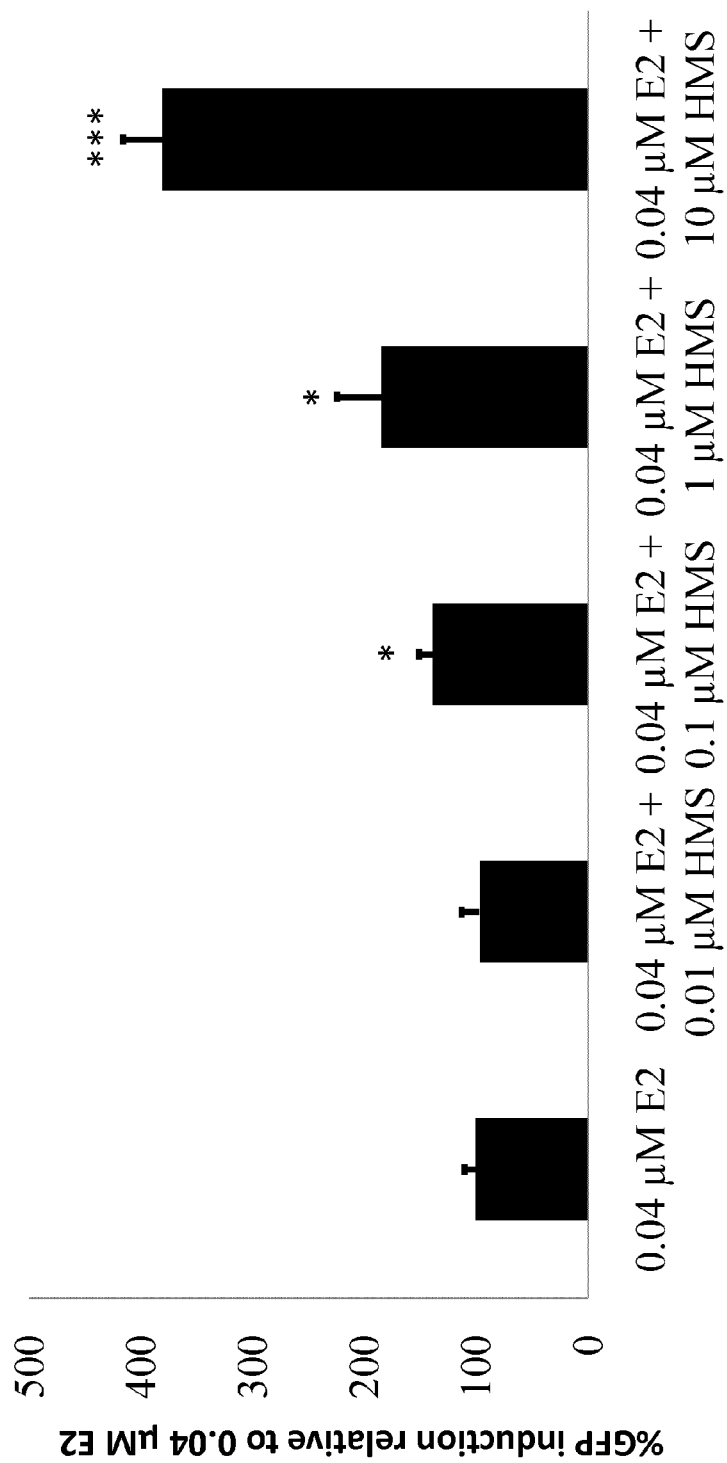


Fig.13

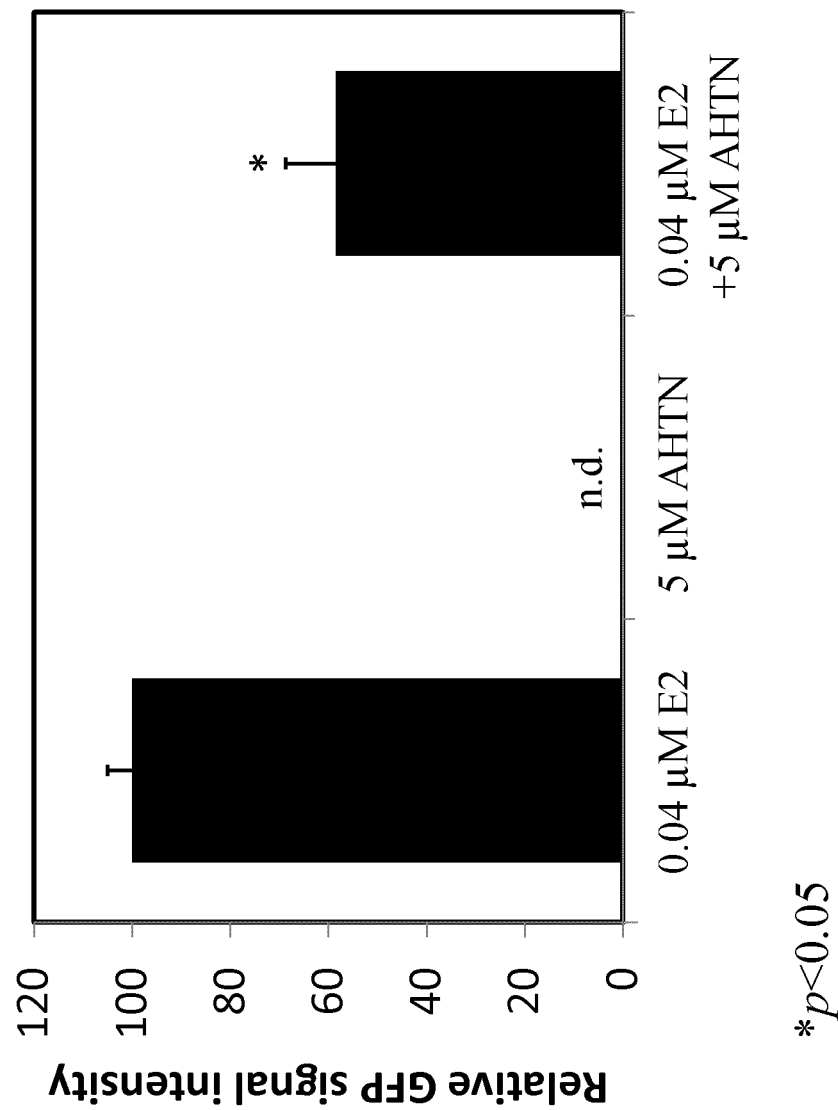


**Fig.14**



\* $p < 0.05$ ; \*\*\* $p < 0.01$

Fig.15





**TRANSGENIC FISH AND USES THEREOF****CROSS REFERENCE TO RELATED APPLICATIONS**

This application is a divisional of U.S. patent application Ser. No. 12/730,956, filed Mar. 24, 2010, the disclosure of which is hereby incorporated herein by reference in its entirety.

**FIELD OF THE INVENTION**

The present invention relates generally to transgenic fish and their use in, inter alia, detecting estrogenic and anti-estrogenic compounds, monitoring estrogen-like activity in the environment, and elucidating liver regeneration.

**BACKGROUND OF THE INVENTION**

Many chemical substances that have been regarded as safe are found to have the activity to disrupt the endocrine system and cause adverse effects in organisms. The wide presence of these kinds of chemicals in the environment has raised attention and concern worldwide. Because of the increasing reports of the estrogenic effects of these substances on male organisms, most of the research efforts have been directed to estrogen-like endocrine disruptors.

It is important to have a rapid, sensitive, economical, and quantifiable method to identify chemical substances that can mimic estrogen or interfere with the estrogen endocrine system, and that can assist in evaluating the biological effects of these chemical substances on organisms. Of the current available methods, in vitro cell line assays and in vivo transgenic fish assays have been investigated. Cell line assays can be rapid and sensitive, but they cannot provide information on toxicodynamic and toxicokinetic effects of test substances on organisms. Currently, transgenic fish assays can overcome some of the deficiencies of the cell line assays. However, current transgenic fish assays are limited in that they cannot be used to monitor endocrine disruptors in freshwater species, and they cannot be used in a marine environment.

**SUMMARY OF THE INVENTION**

One aspect of the present invention relates to an expression cassette that includes: (i) a reporter nucleotide sequence encoding a reporter protein; (ii) an estrogen-responsive 5'-regulatory nucleotide sequence isolated from a medaka fish, where the regulatory nucleotide sequence is operably linked 5' to the reporter nucleotide sequence to induce expression of the reporter protein in the presence of estrogen or an estrogen-like compound; and (iii) a 3'-regulatory region operably linked 3' to the reporter nucleotide sequence.

Another aspect of the present invention relates to a transformation vector that includes an expression cassette of the present invention.

Another aspect of the present invention relates to a host cell transduced with an expression cassette of the present invention.

Another aspect of the present invention relates to a transgenic cell of an aquatic animal, where the transgenic cell includes at least one expression cassette of the present invention.

Another aspect of the present invention relates to a transgenic fish that includes at least one expression cassette of the present invention. The transgenic fish exhibits an observable mark when exposed to a liquid medium containing an estro-

gen or an estrogen-like compound. The observable mark includes a reporter protein expressed by the at least one expression cassette.

Another aspect of the present invention relates to a method of preparing a transgenic fish for use in detecting the presence of an estrogen or estrogen-like compound in a liquid medium. This method involves transforming a non-transgenic fish with the expression cassette of the present invention, thereby yielding a transgenic fish that includes at least one of the expression cassettes.

Another aspect of the present invention relates to a method of screening a liquid medium for an estrogenic substance.

Another aspect of the present invention relates to a method of screening a liquid medium for an estrogenic substance. This method involves exposing a transgenic fish of the present invention to a liquid medium to be tested (i.e., to be tested for the presence of an estrogenic substance). After this exposing step, the method then involves determining whether or not the transgenic fish exhibits an observable mark produced by the induced expression of the reporter gene (contained in the transgenic fish). The presence of the observable mark indicates that the liquid medium contains an estrogenic substance.

Another aspect of the present invention relates to a method of screening for a compound having anti-estrogenic activity. This method involves the following steps: (a) providing a first transgenic fish and a second transgenic fish of the present invention, where the first and second transgenic fish are of the same species and at substantially the same developmental stage; (b) exposing the first transgenic fish to a first liquid medium, where the first liquid medium includes an estrogen or estrogen-like compound; (c) exposing the second transgenic fish to a second liquid medium, where the second liquid medium includes the first liquid medium and a test compound; and (d) comparing the quantified intensity of any observable mark exhibited by the first transgenic fish with the quantified intensity of any observable mark exhibited by the second transgenic fish, where a decrease in the quantified intensity of the observable mark in the second transgenic fish compared to that of the first transgenic fish indicates that the test compound has anti-estrogenic activity.

Another aspect of the present invention relates to a method for investigating the effect of an estrogenic compound on liver regeneration. Generally, this method involves performing a partial hepatectomy on an adult transgenic fish of the present invention, where the partial hepatectomy is effective to remove a portion of the liver of the transgenic fish; exposing the transgenic fish to a test liquid medium containing a test estrogenic compound; and analyzing the liver of the transgenic fish to detect any regeneration of liver tissue.

Another aspect of the present invention relates to a method for investigating the effect of different estrogenic compounds on liver regeneration, as opposed to investigating a single estrogenic compound. Generally, this method involves the following steps: (a) providing a first transgenic fish and a second transgenic fish of the present invention, where the first and second transgenic fish are adults of the same species; (b) exposing the first transgenic fish to a first liquid medium that includes a first test estrogenic compound solution; (c) exposing the second transgenic fish to a second liquid medium comprising a second test estrogenic compound solution; (d) quantifying the intensity of any observable mark exhibited by the first transgenic fish and the second transgenic fish; (e) performing a partial hepatectomy on the first and second transgenic fish, where the partial hepatectomy is effective to remove a portion of the liver of the first and second transgenic fish; (f) repeating steps (b) through (d) of this method; and (g)

analyzing the liver of the first and second transgenic fish to compare the effects of the estrogenic compounds contained in the first liquid medium and the second liquid medium on any regeneration of liver tissue.

#### BRIEF DESCRIPTION OF THE DRAWINGS

The patent or application file contains at least one drawing executed in color. Copies of this patent or patent application publication with color drawings will be provided by the U.S. Patent and Trademark Office upon request and payment of the necessary fee.

FIG. 1 depicts the relative promoter activity of different lengths of omChgH5'-upstream region represented by the frequency of GFP positive larva derived from embryos injected with nucleic acid sequences including the omChgH promoter-EGFP (enhanced green fluorescence protein) in response to 17 $\beta$ -estradiol.

FIG. 2 depicts the effects of mRNA polyadenylation signal of different genes on the relative promoter activity of nucleic acid omChgH promoter-EGFP represented by the frequency of GFP positive larva derived from embryos injected with nucleic acid comprising the omChgH promoter-EGFP (enhanced green fluorescence protein) flanked by olChgH, SV40 or omChgH 3'-flanking region in response to 17 $\beta$ -estradiol.

FIGS. 3A-3G depict the native and induced expression of transgene GFP in transgenic fish produced by introduction of a nucleic acid sequence including the omChgH promoter-EGFP transgene. The expression of GFP is exclusively restricted to the liver of adult female fish (FIG. 3C), but not embryo (FIG. 3A), larva (FIG. 3B), or male fish (FIG. 3D). FIGS. 3E-3G show that the expression of transgene GFP can also be induced in the liver of embryo, larva and male fish by 17 $\beta$ -estradiol.

FIG. 4 depicts a flowchart of one embodiment of the use of transgenic fish larvae of the present invention to analyze the estrogen-like activity of a test solution.

FIG. 5 depicts the estrogen sensitivity of transgenic fish larvae of varying days after fertilization.

FIG. 6 is a graph showing that pH ranging from 5 to 9 does not affect the response of transgenic fish larvae to 17 $\beta$ -estradiol.

FIG. 7 is a graph showing that salinity ranging from 0 to 35 ppt does not affect the response of transgenic fish larvae to 17 $\beta$ -estradiol.

FIG. 8 depicts a time-lapse increase of GFP signal intensity in transgenic larvae in response to estrogen.

FIG. 9 depicts a dose-response of transgenic fish larvae to 17 $\beta$ -estradiol.

FIG. 10 depicts the dose-response of transgenic fish larvae to synthetic hormone 17 $\alpha$ -ethinylestradiol.

FIG. 11 depicts the liver GFP signal induced by phytoestrogens genistein, genistin, daidzein, and daidzin separately and their combination extracted from soy milk.

FIG. 12 depicts the liver GFP signal induced by UV filters BP-3 and 4-MBC separately and in combination. Transgenic fish larvae of one embodiment of the present invention are exposed to 4-MBC (5  $\mu$ M), HMS (5  $\mu$ M) and the combination of these two compounds mixed at their half concentrations (2.5  $\mu$ M). The GFP signal induced by their mixture is significantly higher than that of these two compounds.

FIG. 13 depicts the liver GFP signal induced by 17 $\beta$ -estradiol alone or in combination with UV filters B-MDM, OMC or OD-PABA.

FIG. 14 shows that the liver GFP signal induced by 17 $\beta$ -estradiol can be increased by HMS in a dose-dependent manner.

FIG. 15 shows that the liver GFP signal induced by 17 $\beta$ -estradiol can be decreased by polycyclic musk AHTN.

#### DETAILED DESCRIPTION OF THE INVENTION

In one aspect, the present invention relates to a transgenic fish that contains an expression cassette that includes an estrogen-responsive 5'-regulatory nucleotide sequence isolated from a medaka fish, where the estrogen-responsive 5'-regulatory nucleotide sequence is operably linked 5' to a reporter nucleotide sequence. When the transgenic fish is exposed to a liquid medium containing estrogen or an estrogen-like compound, the reporter nucleotide sequence expresses a corresponding reporter protein that can be detected in vivo or in vitro. The transgenic fish of the present invention is useful for a variety of applications, including, without limitation, for use in detecting estrogenic compounds and anti-estrogenic compounds, monitoring estrogen-like activity in the environment, and elucidating liver regeneration.

The transgenic fish of the present invention can be of various freshwater, brackish water, or saltwater (marine water) species of fish, including, without limitation, fish of the *Oryzias* species and the *Danio* species. Fish in the *Oryzias* genus belong to the Adrianichthyidae family and include, for example, *Oryzias melastigma* (alternative name *Oryzias dancena*) (Marine or brackish medaka), *Oryzias latipes* (Japanese medaka), *Oryzias celebensis*, *Oryzias marmoratus*, *Oryzias matanensis*, *Oryzias nigrimas* (black buntingi), *Oryzias orthognathus* (buntingi), and *Oryzias profundicola*. Fish in the *Danio* genus belong to the Cyprinidae family and include, for example, *Danio rerio* (zebra fish), *Danio albolineatus*, *Danio abolineatus*, *Danio choprae*, *Danio dangila*, *Danio erythromicron*, *Danio feegradei*, *Danio kerri*, *Danio kyathit*, *Danio margaritatus*, *Danio meghalayensis*, *Danio nigrofasciatus*, and *Danio roseus*.

With regard to the brackish medaka fish (*Oryzias melastigma*), this species is one of many medaka species identified as belonging to the *Oryzias* genus (some species being set forth herein above). The brackish medaka fish is native to coastal waters and fresh waters in Pakistan, India, Burma and Thailand (K. Naruse, *Fish Biol. L. Medaka* 8:1-9 (1996)), and thrives in waters of varying salinity ranging from 0 parts per thousand (ppt) to as high as 35 ppt. Additionally, this brackish medaka fish has a number advantages for transgenic development, including: (1) small size (2-3 cm for adult fish); (2) relatively short generation time (2-3 months); (3) dimorphic sex (e.g., females have a flat distal surface of the anal fin, while that of males is convex due to separated longer fin rays); (4) high prolific capacity to reproduce; (5) translucent eggs and larvae (up to 15 days post fertilization), which facilitates the positioning of DNA microinjection needles and observation of internal organs; and (6) adaptable to various transgenic techniques used to produce transgenic fish of other *Oryzias* species (e.g., *Oryzias latipes*).

Regarding the highly prolific capacity of the brackish medaka fish to reproduce, spawning of this fish can be induced all year round, and each pair of female and male fish can produce 20-30 eggs daily for up to several months under indoor maintained conditions (e.g., 28 $\pm$ 1 $^{\circ}$  C. with a constant light cycle of 14 h-light/8 h-dark and fed with commercial hormone-free flake food and brine shrimp (*Artemia salina*)). Eggs usually hatch in 11 to 15 days at 28 $\pm$ 1 $^{\circ}$  C.

*Oryzias latipes* has been used to produce transgenic fish carrying a variety of transgenes (e.g., K. Ozato et al., *Cell Differ.* 19:237-244 (1986); K. Inoue et al., *Cell Differ. Dev.* 27(1):57-68 (1989); E. Tamiya et al., *Nucleic Acids Res.* 18:1072 (1990); K. Inoue et al., *Cell Differ. Dev.* 29(2):123-

128 (1990); J. Lu et al., *Mol. Marine. Biol. and Biotechnol.* 1(4/5):366-375 (1992); H. J. Tsai et al., *Mol. Mar. Biol. Biotechnol.* 4(1):1-9 (1995); R. N. Winn et al., *Marine Env. Res.* 40(3):247-265 (1995)). Studies have shown that the two medaka species of *Oryzias melastigma* and *Oryzias latipes* share high morphological, physiology, and genomic similarity, and that it is easy to adapt the transgenic technique from *Oryzias latipes* to the brackish medaka *Oryzias melastigma* (X. Chen et al., *Ectotoxicol. Environ. Saf* 71:200-208 (2008); X. Chen et al., *Comp. Biochem. Physiol. C. Toxicol. Pharmacol.* 149:647-655 (2009)).

In one embodiment, the transgenic fish of the present invention is engineered to contain one or more copy of a linearized nucleic acid sequence containing the promoter region of an estrogen target gene, which is operably ligated to the coding region of a reporter gene (e.g., a fluorescence protein gene), and then flanked by a termination sequence (e.g., an mRNA polyadenylation signal sequence). The pro-

motor region sequence and the termination sequence can originate from the same estrogen target gene, although the present invention does not require that they have the same origin.

A suitable promoter region can be one that is derived from an estrogen-dependent gene. In one embodiment, a particular estrogen-dependent gene contemplated by the present invention can include, without limitation, the choriogenin H genes from *Oryzias melastigma* (the gene being abbreviated as "omChgH") and *Oryzias latipes* (the gene being abbreviated as "olChgH").

The omChgH gene of *Oryzias melastigma* has been characterized as being highly sensitive to estrogen and estrogen-like substances. The complete coding sequence of the omChgH gene of *Oryzias melastigma* has been determined and deposited as GenBank Accession No. EF392365. As used herein, the omChgH gene coding sequence corresponds to SEQ ID NO:1, as follows:

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1  gaattcacta gtgattacta tagggcaccg gctggtcgacg gcccgggctg gtatcgtgag
61  ccatgtatcg cgtatcgtat cgtatcgtga ggtaccacgt gattcccacg cctaataatt
121 atggcctaaat actatatatg tagaaatcta actgttgtta aaaaagccag agttttttta
181 atcttcacaa agaaattggt ttgcaaatta atgaaaatcc aatgcaaaag ctgttaggac
241 agtcgaagcc tggacttggt tggcatcata ttttattatt attattatga ttcctttctt
301 ttgttaccac ctggcaaggt catatcaatt attctgtaat tctcggtatt ggatcattta
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481 tgtagtgata aagctccaaa tttattttgt ttattttccc aaatccatca ttatttcaaa
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1921 tgttacttat tatctcatga aatcctttgt actgttttgc cttgggggtt ttattgacta  
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2701 gtcaaaaaa gtgacacagt tggccagatc taaagtctcg gatgttgtgt tctggtcaaa  
2761 taaacatatg acaatgaata aaagttaggg tcacatggac attttgaaaa gggcttttat  
2821 gacctagagt ttgatttgta gataaacccc tttatgtcaa actgtcatca aacacaaccc  
2881 tgatccaaca gcgttacccc ttagctatca cagttcactt tgtaaatgtg tcacaaaaaa  
2941 tgggtcataa tcatcaaaat gaattataat ttctaagaaa ttatagcatt tagatattgc  
3001 acctttttta acacctcaaa attactgggt tcttttcttt ttattatttc tttagtttta  
3061 agtttgtttt gaaaatgac ttaattattgc agcggagtag cacgtattta actaaaatta  
3121 aaactactaa ccatgaacat ttttcagcat ttgtgtgggt tagtttttgt gacatgttg  
3181 catgtagtca tattttgagt ttttgaggtg ctttttaaca gctttcagaa gcagtttaac  
3241 catttaaaat taagaaaatt caagtcaga ttacactttt catattatct tagtttgtga  
3301 aaaaatagta ttataatata ttagaagact ttcttctaca caggactacc tgaggtttaa  
3361 acaacacttt atcaactgag aggtagcact gaaatatgtg cgtgcaagt tatttaattg  
3421 tttttttttt ttattttttt agctcttget acctagaaac attgtaaaaa aaattactga  
3481 aactttacac attaattttc tgaagacctc atcataatcc atgatgttta acgggtttga  
3541 attgcaacta tttttactgc atcataatta ccacatccac tctattaaat tctattccaa  
3601 taccagagtt aaggctagag ttgtgtacac tacgaccgat ccacagttat gcaacgtaat  
3661 tctgagaaac tatgagtctc ctttattatt cctctatcaa accagtgggt aaggaagtat  
3721 ttttaatttt acttaaaaaa aacttttcaa cattaaacaa ttcaaagtgt tgtgcgtctg  
3781 tctgaattca ttttaattat cgttaattga tttctacac aattaatact tgcatatgtt  
3841 ttgctttttt taaatgctaa ctttattaca tttctgatt ggggctaaaa atcaactgga  
3901 aagaaaatgg ttttttttac acttcattat gtcctgtttg gggttgctgg agcctgtccg  
3961 agttacctca ggggaaaagc cagagtgcac cctgaacatt gagaggtcac acacagacac  
4021 atccatccat gttcttaacc tgctaaatcc cttattaagc atgattttgg actgtctgaa  
4081 gaaaagccaa gcaagaacat gccaaagcca cgcaaaaggg tcccaacat gctttcaacc  
4141 aggaccagct tactgtgagg tgagaccgct aactactgca ccacagtga gcccatat

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4201 agacttaatg caatatattg tgttttggg aagaagctgg agagcctgga gggaaccagt  
4261 gcatggggaa aacatacaaa cctcacacag caaggatcac atctgcctca gtatgtattt  
4321 ttacagctta ctgtaattaa aaacagtaaa aaaaaaaaaa ctaaaaaaaa aaacatcaaa  
4381 ttgccacaaa aagtacatga tttaaaaaaa ggtttctgtc tacatgttta catcttaatt  
4441 tgatttaaaa cactaaaaaa tacataagat tgcccccata aaatgcaatt ataaaaagca  
4501 catgatttaa aaaaaaaaaa gtatccgtac gtatttttag atcttaattt ggcttaaaaa  
4561 ctaaaaaata cattagattg ccaaaaaatg caattataaa tatacatgtt ttttaataaa  
4621 agccatcagc tgcacatgca ttttgggaaa aaaaggttgt cttatagaaa aagtcacat  
4681 ccattacaca atttaactgt ttactttgaa taaaaacaca ttactgtgct gctttattat  
4741 tgctctcact ctatggggtt caaactaatg tgtatttgct atttttgtgc tttgtttagt  
4801 ggatatctaa cagtattgca acagactgaa agggcaaaaag aggtaattac ttcttaaaaa  
4861 gctaacatta gattgtttct gtcattgaaa atgcatgaaa acatactgca acaatttatt  
4921 ttaatgttca ttaagtgtca aggttttttag gattcttgaa atcctcaaat tgatatgtta  
4981 gcgacacata tttgactata atgggtctct gttatttcat tcatttatgt aacaaaaaca  
5041 ataacacaaa aatgtacata acttttcatt tactcgagac ctattagtta aaaaaatgtg  
5101 gaggaatgag ctttttcttt actacgtcga aatataaaat ttctgagaca taaatcaatt  
5161 ataaatatag agtatatggc ctgattaaaa aaaagaaaaa aacttttcaa ctgattttgt  
5221 aatgcagaaa atcatgctta gcacaaccag agcattcgcc aacataata tttgatgtgg  
5281 acatcatcct aataacctca tataaaagggt tttttttgac caaaatgtgt acaatatgag  
5341 gttttgttgc agattcaggg aaaagatcac tttgtttgtc attgcctact atcaaaacaa  
5401 acatttgagg acagataagt tcgagactgc aaagaccctg aaaaggtctc catgacctgg  
5461 atgtcacaaa agcctttcat tcattccaac gcaacgacct gatctggcat ttcacgcaaa  
5521 ggacagaata gtccacatga attacataaa attgacttaa caaaacacac cctgaagcgt  
5581 ttgtgattgg gagctacttg gtttgagagg tggagtttga agagcatcaa aggtaaagac  
5641 acataaatag agcaggagag ggaaatttac acttagggac ccacggtgac agacagctgt  
5701 gggaccatgg caaggcactg gagtattacg gttttttccg cactagctct gatatgttct  
5761 ttcttgoga cccaagtgga tgctcagaaa ggccccctc aagaccctaa ggttccatc  
5821 cctccatct atccacagcc gaagccgag gacctcaac acgtttcacc gccttacaac  
5881 ccaggaagc cgcagtatcc aggaagcca cagtatccag ggaagccgca agtccacag  
5941 tatcctcaga cccctcagta tcctcagacc cctcagcagc cgcagagtcc acagtatcct  
6001 cagaccctc agtatcctca gaccctcag cagccgaga gtccctcagta tcctcagtcc  
6061 cctcagtatc ctcagactcc tcagtccct cagtatctc agtccctca gtatcctcag  
6121 actcctcagt atcctcagaa tcctaagggtg tatgggtgatg acagttctaa gccttcaact  
6181 ccgtcaaaag ctagctatcc tcagcctcag gccccccagt acccatctaa gcctcaagct  
6241 cccagctgc ctcaggcccc ccagtacca actaagcctc aagctccca gctgcctcaa  
6301 gctccccagc tgctcaggc cccccagtac ccaactaagc ctcaagctcc tcagtatcct  
6361 caagctcctc agcagcctca ggccccccag tacccaacta agcctcaagc tcctcagcag  
6421 cccaggtctc ccagtagccc aactaagcct caagctcctc agtaccctca agctcctcag  
6481 cagccccagg ctcccagta cccaacaaaa cctcagcagc cccagtacc aacaaagcct  
6541 cagtctcctc agtaccctca agatcctaaa aatccaaatc ctcagaatcc tcctcttcat

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6601 cctccccctg ttaagagctg tgaggtgcc cgcgatgtga gagtcccatg tggagttcca  
 6661 gacatctctc cttctgcatg cgatgccatt gactgctgtc atgatggcca agcctgctac  
 6721 tttggaacag gaggttaagt gtttctccag ctgctatgat cagaggcttt ttgtaagggtg  
 6781 acggctgatc gtgcaatcgt cgatcccatc tatttcttct agcaaccgtt cagtgcacca  
 6841 aggacggaca cttcatcggt gtggtggcca aggatgtcac cttaccccat ctcgatcttg  
 6901 aaactatttc acttttggga ccgggtcaag aatgtggagc tgttgactct aattcagctt  
 6961 tcgccatcta ctactttccc gtcaactcagt gcggcactca tgtcacggta acactcagtc  
 7021 ttgtttatat cttatagtca taaggtcaat ctttgagatt ctatccttct tattgttaaa  
 7081 ttttgaacca ttaaaaggaa gagcccgagg ttatagtcta tgaaaaccgg atgacatcct  
 7141 catatgaagt tggagttgga ccgcttgagg ctattaccag agacagctct tttgagtagg  
 7201 tcatcatttg tgtttagat caaacagatt tactaatgtc taactaatat ctatcagggg  
 7261 taaacagaat catgcacagt gtattaacac agttcttttt ctcaggctcc tcttccagtg  
 7321 cagataccat gcaacatctg ttgaaactct ggtcgtggaa gtgctgccag tggatagtc  
 7381 tctttccatt gctgagcttg gacccctcaa tgtgtacttg caaattgcc aaggagtagg  
 7441 tcagacaaag ggctgcgacg aaggtcagtg cacggcagtc tggcacagcc agtgtgttg  
 7501 tcaaacattt gaaaaagctg cctgtcgtaa cgtttgtttg ctcccacagt ggccgcagcc  
 7561 tacacctctt tctacacgga tgccgactat cctgtgacca aagtactgag agaaccagtc  
 7621 tatgtggacg ttcaaatcct tggcagaaca gatccaaatc tggttctgac tcttggacgc  
 7681 tgttgggcaa ccacaagccc caatgctttc agtctgcccc agtgggacat tttgattgac  
 7741 gggtaaaaaa aaaaaatct accaattcat tccataaaga ccatttttgt tcaaaactaag  
 7801 ctccaaatct cacacttttt agccgaatta ctaaatatct aaccaaccat tacttcttct  
 7861 ttaccttttt tccatccaga tgtccatatg aagatgatcg ctacctgtct gcgttggttc  
 7921 caatcgattc ctctctggtt ttgccattcc caactcatca cagacgcttc ttattcaaga  
 7981 tgtttacott tgttgatcct cattcaatgg aacctactaag ggaagagggt ggtactgaat  
 8041 tactcaagta gaggtttaac ttggttctta acctgtactt ttctttcagg tgtacattca  
 8101 ctgcagcaca gctgcatgcg ttccaggaca gggtagcagt tgtgaacct catgcagcag  
 8161 aagaagtagg gggtctattt ataaccgttc acatgttttt ttgtgtgttg taatgacct  
 8221 gtccactcat cattggccat gttttgtgtc tttttgtaga aggaagagat actgacgctg  
 8281 tatccattag aacggatgaa agaaagggtg tggatcgtc tggagaagt ctcattggtg  
 8341 ccgaagctgc tggacagtct taactgtgaa ccgacagaag ctccagagtt cggaaaaaat  
 8401 aacataacat tatgaaaatc tgtttcatca tggttcagaa ttaaatgcat aaagt

The estrogen-responsive 5'-regulatory nucleotide sequence of the present invention can include, without limitation, any sequence of the promoter region of the omChgH gene of SEQ ID NO:1.

In one embodiment, a suitable estrogen-responsive 5'-regulatory nucleotide sequence of the present invention comprises a nucleotide sequence corresponding to SEQ ID NO:2, as follows:

ctcgtagctc caaaacccaa catattttgt ctataaagg ttagcttaaa aatatatatt	60
ttttcaatta aaagaaaata actgaaaaac tggaaattta tcttaaaact accaagaagt	120
tatctattta atcattatgt ttacattcaa gtcgtcttag ggccacaagg tacttagcct	180
ttgctgttat ttacaaagac gacaatttaa gtgtatgaag tgatctttat aaatcagtc	240
aatatatatt tttaaatgca ctcatcctcc ttggcagca agttgtggcg aaaatccttt	300
ataatcttca aattacaagt atgaacaata ttgtttcctt taaaacaatt agcagtctaa	360

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tcatttttcta taatatgaaa gttcatatct tatgtagttg aaaagtggtgc tatttttcta	420
attgtatttc tttttatata ttttgcatct agtgaactac tgcattaact agcttgcaat	480
acttttcgca ttgtgtctct cataaaaaat atagatatgg ttatttataa aaatgcgttt	540
actcttaatt gttgcctttt ggagtaaaat tttatgcttt tgtctgtatt gcctttttga	600
cattcagaag aggatgcttc tgtatgacgg cgtattaagg ccttgagggt atcatgtctg	660
tggtgtgact gaatgattac attgatgtg ttttacctt tgaaaatcac catttgaaac	720
caagttttgt ctcacaattg catcaaacag aaaaaaac atcaaagcaa gctaattgaa	780
tgttgattgg tttaatccaa gtcattacag tttattccac ctgagtctag atgattctaa	840
cagattgttt gactgtgtgc aaaatgatcc tacatgaatg caagtccaaa gcaagataaa	900
tgctctttct cttcaaatga ttgacatta atctttccct aaaatttatt attggtcata	960
atattgataa atctttgttt cgtgatgcat gagaccttat cacctaattt tgcagtaaag	1020
agagtcaact aaaaaatttt ctgacactct tttgttactt attatctcat gaaatccttt	1080
gtactgtttt gccttggggg ttttattgac taacatctct ttaatcacac ataattcaac	1140
tgttcataat tccaaatag ttttagcttt tgttataacg atttaagtta cagcaatgtg	1200
ttttattttt ttctgtattt aactgaacta tgttgccctt ataaacagtt acatgtatgt	1260
aataggagtg gcctgatatt gccacacacc cctcatcgga tgctatgtca tggacaaata	1320
gctccactat atggtc aaac ataagtataa cttaagaaag taacagaaaa atttaattta	1380
aaaaaagttt tttttttttt ttgtcttttc catttcttta ttttatttat ttaaatacaa	1440
aaaaagtc aa atcctttact tctcggcaca taatttttca aacaatcttt ttaaataaaa	1500
ttattaatta aagcaaatat tcattcacaa aaaaaactt ttagagtttg aaaaatgcga	1560
atttaaggtt gtattattat aatcaatatt gatttaagtt tacaataga tttaaaaaaa	1620
aaatggaaat ctgctatgtt tccctgacag ccttcaaaag tttccttctg gaaaagttaa	1680
actgttgttt tgtgacgggt ttttttttta aacgctttgt ctggatcaat taaagcatag	1740
ttttttcaca tttgcggttt ttcacggaga gaggaagttt cctttattgt gtatcctgcc	1800
ctgagtacca gtgacattag agcctgtaac gtgtcaaata aagtgcaca gttggccaga	1860
tctaaagtct cggatgttgt gttctggtca aataaacata tgacaatgaa taaaagttag	1920
ggtcacatgg acattttgaa aagggtttt atgacctaga gtttgatttg tagataaacc	1980
cctttatgtc aaactgtcat caaacacac cctgatccaa cagcgttacc ccttagctat	2040
cacagttcac tttgtaaat tgtcacaaaa aatgggtcat aatcatcaaa atgaattata	2100
atttctaaga aattatagca tttagatatt gcacctttt aaacacctca aaattactgg	2160
tttcttttct ttttattatt tctttagttt taagtttggt ttgaaaatga tcttaattat	2220
gcagcggagt agcacgtatt taactaaaat taaaactact aacctgaac atttttcagc	2280
atttgtgtgg tttagttttt gtgacatgtt tgcattgtat catattttga gtttttgagg	2340
tgctttttta cagctttcag aagcagttta accatttaaa attaagaaaa ttcaagctca	2400
gattacactt ttcattattat cttagtttgt gaaaaatag tattataata tattagaaga	2460
ctttcttcta cacaggacta cctgaggttt aaacaacact ttatcaactg agaggtagca	2520
ctgaaatatg tgcgctgcaa gttatttaat tgtttttttt tttttatttt ttagctcttg	2580
ctacctagaa acattgtaaa aaaaattact gaaactttac acattaattt tctgaagacc	2640

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tcatcataat ccatgatggt taacggggtt gaattgcact tatttttact gcatcataat	2700
taccacatcc actctattaa attctattcc aataccagag ttaaggctag agtttggtac	2760
actacgacgg atccacaggt atgcaacgta attctgagaa actatgagtc tcctttatta	2820
ttcctctatc aaaccagtggt tgaaggaagt attttaattt ttacttaaat aaaacttttc	2880
aacattaaac aattcaaatg tgtgtgcgtc tgtctgaatt catttaatta ttcgttaatt	2940
gattttctac acaattaata ctgcatatg ttttgccttt tttaaagtct aactttatta	3000
cattttctga ttggggctaa aaatcaactg gaaagaaaat gggttttttt acacttcatt	3060
atgtcctggt tgggggtgct ggagcctgct cgagttacct caggggaaaa gccagagtgc	3120
acctgaaca ttgagaggtc acacacagac acatccatcc atgttcttaa cctgctaaat	3180
cccttattaa gcatgatttt ggactgtctg aagaaaagcc aagcaagaac atgccaagtc	3240
cacgcaaaag ggtcccaacc atgctttcaa ccaggaccag cttactgtga ggtgagaccg	3300
ctaactactg caccacagtg cagcccatat ttagacttaa tgcaatatat tgtgttttgg	3360
ggaagaagct ggagagcctg gagggaacca gtgcatgggg aaacatata aacctcacac	3420
agcaaggatc acatctgcct cagtatgtat ttttacagct tactgtaatt aaaaacagta	3480
aaaaaaaa aactaaaaaa aaaaacatca aattgccaca aaaagtacat gatttaaaaa	3540
aagggttctg tctacatggt tacatcttaa tttgatttaa aactactaaa aatacataag	3600
attgccccca aaaaatgcaa ttataaaaag cacatgattt aaaaaaaaa aggtatccgt	3660
acgtattttt agatcttaat ttggcttaaa cactaaaaaa tacattagat tgccaaaaaa	3720
tgcaattata aatatacatg ttttttaata aaagccatca gctgcacatg cattttggga	3780
aaaaaagggt gtcttataga aaaagtcac atccattaca caatttaact gtttactttg	3840
aataaaaaca cattactgtg ctgctttatt attgctctca ctctatgggg ttcaaaactaa	3900
tgtgtatttg tcatttttgt gctttgttta gtggatatct aacagtatgt caacagactg	3960
aaagggcaaa agaggtaatt acttctctaa cagctaact tagattgttt ctgtcatgga	4020
aatgcatga aaacatactg caacaattta ttttaagtgt cattaagtgc taagggtttt	4080
aggattcttg aaatcctcaa attgatatgt tagcgacaca tatttgacta taatggtctt	4140
ctgttatttc attcatttat gtaacaaaaa caataacaca aaaatgtaca taacttttca	4200
tttactcgag acctattagt taaaaaatg tggaggaatg agccttttct ttactacgtc	4260
gaaatataaa atttctgaga cataaatcaa ttataaatat acagtatatg gcctgattaa	4320
aaaaaagaaa aaaacttttc aactgatttt gtaatgcaga aaatcatgct tagcacaacc	4380
agagcattcg ccaacatata catttgatgt ggacatcatc ctaataacct catataaaag	4440
gttttttttg accaaaatgt gtacaatatg aggttttggt gcagattcag ggaaaagatc	4500
actttgtttg tcattgccta ctatcaaaac aaacatttga ggacagataa gttcgagact	4560
gcaaagaccc tgaagggtc tccatgacct ggatgtcaca aaagccttcc attcattcca	4620
acgcaacgac ctgatctggc atttcacgca aaggacagaa tagtccatc gaattacata	4680
aaattgactt aacaaaacac acctgaagc gtttgtgatt gggagctact tggtttgaga	4740
ggtggagttt gaagagcatc aaaggtaaag acacataaat agagcaggag agggaaaattt	4800
acacttaggg acccatcggg tcagacagct gtgggacc	4838

The estrogen-responsive 5'-regulatory nucleotide sequence of SEQ ID NO:2 corresponds to a 4827 base pair region 5' upstream of the omChgH gene of SEQ ID NO:1. 65 Base pairs 4828-4838 of SEQ ID NO:2 correspond to a 5'-untranslated region of the omChgH gene of SEQ ID NO:1.

In one embodiment, a suitable estrogen-responsive 5'-regulatory nucleotide sequence of the present invention comprises a nucleotide sequence corresponding to SEQ ID NO:3, as follows:



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tttaggattc ttgaatcct caaattgata tgtagcgac acatatttga ctataatggt    60
cttctgttat ttcattcatt tatgtaacaa aaacaataac acaaaaatgt acataacttt    120
tcatttactc gagacctatt agttaaaaa atgtggagga atgagccttt tctttactac    180
gtcgaaatat aaaatttctg agacataaat caattataaa tatacagtat atggcctgat    240
taaaaaaaag aaaaaaactt ttcaactgat ttgtaatgc agaaaatcat gcttagcaca    300
accagagcat tcgccaacat atacatttga tgggacatc atcctaataa cctcatataa    360
aaggtttttt ttgacaaaaa tgtgtacaat atgagggtttt gttgcagatt cagggaaaag    420
atcactttgt ttgtcattgc ctactatcaa aacaaacatt tgaggacaga taagttcgag    480
actgcaaaga ccctgaaaag gtctccatga cctggatgtc acaaaagcct ttcattcatt    540
ccaacgcaac gacctgatct ggcatttcac gcaaaggaca gaatagtcca catgaattac    600
ataaaattga cttaacaaaa cacaccctga agcgtttgtg attgggagct acttggtttg    660
agaggtggag tttgaagagc atcaaaggta aagacacata aatagagcag gagagggaaa    720
tttacctta gggacccatc gggtcagaca gctgtgggac c                          761

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The estrogen-responsive 5'-regulatory nucleotide sequence of SEQ ID NO:3 corresponds to a 750 base pair region 5' upstream of the omChgH gene of SEQ ID NO:1. Base pairs 751-761 of SEQ ID NO:3 correspond to a 5'-untranslated region of the omChgH gene of SEQ ID NO:1.

In one embodiment, a suitable estrogen-responsive 5'-regulatory nucleotide sequence of the present invention comprises a nucleotide sequence corresponding to SEQ ID NO:4, as follows:

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gttgcagatt cagggaaaag atcactttgt ttgtcattgc ctactatcaa aacaaacatt    60
tgaggacaga taagttcgag actgcaaaga ccctgaaaag gtctccatga cctggatgtc    120
acaaaagcct ttcattcatt ccaacgcaac gacctgatct ggcatttcac gcaaaggaca    180
gaatagtcca catgaattac ataaaattga cttaacaaaa cacaccctga agcgtttgtg    240
attgggagct acttggtttg agaggtggag tttgaagagc atcaaaggta aagacacata    300
aatagagcag gagagggaaa tttacctta gggacccatc gggtcagaca gctgtgggac    360
c                                361

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The estrogen-responsive 5'-regulatory nucleotide sequence of SEQ ID NO:4 corresponds to a 350 base pair

region 5' upstream of the omChgH gene of SEQ ID NO:1. Base pairs 351-361 of SEQ ID NO:4 correspond to a 5'-untranslated region of the omChgH gene of SEQ ID NO:1.

The 3'-regulatory region of the present invention can include, without limitation, a choriogenin H3'-flanking region that includes choriogenin H 3'-untranslated region (represented as UTR). In one embodiment, a suitable 3'-regulatory region can include the omChgH 3' UTR deposited as

GenBank Accession No. DQ778335, which corresponds to the nucleotide sequence of SEQ ID NO:5, as follows:

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ctgtgaaccg acagaagctc cagagttcgg aaaaaataac ataacattat gaaaatctgt    60
ttcatcatgg ttcagaatta aatgcataaa gtgaaaaatc tgttcagagt gtttggaatg    120
tattgcaaaa acaaaacaag ttgatacata aaggtagcaa catttcttca catttcatgt    180
aaaaaaaaa aaaaagtttc ttcacatcag tatagcaggt gtgtagatag agttgacaaa    240
atacaacact tccaccttaa tattctatat ggatcaaatg tgactgtttt cagtgaaacg    300
tcacgacaat aaagtcacat aatacatttc acttttacac aaatttttac tgtctgtttc    360
tgttcttaaa catacaagca ctgaaaacag agatgaatcc agtataacca aacaactcaa    420
acgacaataa aaaaaacaaa aaaattgttt tattatttta aaatgtttta aaaaagtcca    480
attttttaat caaagtaggt caaccatttt taatactgga tcaacaaaca aaaacaatta    540
acaaaaaaaa tcagagttaa tggaaggtaa acacacacat ccgtgaagac aaaaacacaa    600

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gattattatt taaaaactga actagacagc ttactttctca gaaatctgcg actgtaagga 660  
 aaactgtttt cctgttgct ttcaatttgt aaaattgaaa gatgtcaata aataatttac 720  
 cctcttgcat ttgaaaaaca attgtacttt cttggaagaa tatttggcat aaatgcattg 780  
 ttacatgtgg atgtcggtt ttttaagcacc tgctatggtg agcttgggcc a 831

The practice of the present invention will employ, unless otherwise indicated, conventional techniques of chemistry, molecular biology, microbiology, recombinant DNA, and immunology, which are within the capabilities of a person of ordinary skill in the art. Such techniques are explained in the literature. See, e.g., J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, *Molecular Cloning: A Laboratory Manual, Second Edition, Books 1-3*, Cold Spring Harbor Laboratory Press; Ausubel, F. M. et al. (1995 and periodic supplements) *Current Protocols in Molecular Biology*, ch. 9, 13, and 16, John Wiley & Sons, New York, N.Y.); B. Roe, J. Crabtree, and A. Kahn, 1996, *DNA Isolation and Sequencing: Essential Techniques*, John Wiley & Sons J. M. Polak and James O'D. McGee, 1990, *In Situ Hybridization: Principles and Practice*; Oxford University Press; M. J. Gait (Editor), 1984, *Oligonucleotide Synthesis: A Practical Approach*, Irl Press; D. M. J. Lilley and J. E. Dahlberg, 1992, *Methods of Enzymology: DNA Structure Part A: Synthesis and Physical Analysis of DNA Methods in Enzymology*, Academic Press; Using Antibodies: A Laboratory Manual: Portable Protocol NO. 1 by Edward Harlow, David Lane, Ed Harlow (1999, Cold Spring Harbor Laboratory Press, ISBN 0-87969-544-7); Antibodies: A Laboratory Manual by Ed Harlow (Editor), David Lane (Editor) (1988, Cold Spring Harbor Laboratory Press, ISBN 0-87969-3, 4-2), 1855. Handbook of Drug Screening, edited by Ramakrishna Seethala, Prabhavathi B. Fernandes (2001, New York, N.Y., Marcel Dekker, ISBN 0-8247-0562-9); and Lab Ref: A Handbook of Recipes, Reagents, and Other Reference Tools for Use at the Bench, Edited Jane Roskams and Linda Rodgers, 2002, Cold Spring Harbor Laboratory, ISBN 0-87969-630-3. Each of these general texts is herein incorporated by reference.

The term "transformation" refers to the transfer of a nucleic acid fragment into the genome of a host cell, resulting in genetically stable inheritance. A "host cell" is a cell that has been transformed, or is capable of transformation, by an exogenous nucleic acid molecule. Host cells containing the transformed nucleic acid fragments are referred to as "transgenic" cells, and organisms comprising transgenic cells are referred to as "transgenic organisms."

"Transformed," "transduced," "transgenic," and "recombinant" refer to a host cell or organism into which a heterologous nucleic acid molecule has been introduced. The nucleic acid molecule can be stably integrated into the genome generally known in the art and are disclosed in Sambrook and Russell, *infra*. See also Innis et al. (1995); and Gelfand (1995); and Innis and Gelfand (1999). For example, "transformed," "transformant," and "transgenic" cells have been through the transformation process and contain a foreign gene integrated into their chromosome. The term "untransformed" refers to normal cells that have not been through the transformation process.

A "transgenic" organism is an organism having one or more cells that contain an expression vector.

"Genetically altered cells" denotes cells which have been modified by the introduction of recombinant or heterologous nucleic acids (e.g., one or more DNA constructs or their RNA

counterparts) and further includes the progeny of such cells which retain part or all of such genetic modification.

As used herein, the terms "polynucleotide," "nucleotide," and "nucleic acid" are intended to be synonymous with each other. "Polynucleotide" generally refers to any polyribonucleotide or polydeoxyribonucleotide, which may be unmodified RNA or DNA or modified RNA or DNA. "Polynucleotides" include, without limitation, single- and double-stranded DNA, DNA that is a mixture of single- and double-stranded regions, single- and double-stranded RNA, and RNA that is a mixture of single- and double-stranded regions, hybrid molecules comprising DNA and RNA that may be single-stranded or, more typically, double-stranded or a mixture of single- and double-stranded regions. In addition, "polynucleotide" refers to triple-stranded regions comprising RNA or DNA or both RNA and DNA. The term polynucleotide also includes DNAs or RNAs containing one or more modified bases and DNAs or RNAs with backbones modified for stability or for other reasons. "Modified" bases include, for example, tritylated bases and unusual bases such as inosine. A variety of modifications can be made to DNA and RNA; thus, "polynucleotide" embraces chemically, enzymatically, or metabolically modified forms of polynucleotides as typically found in nature, as well as the chemical forms of DNA and RNA characteristic of viruses and cells. "Polynucleotide" also embraces relatively short polynucleotides, often referred to as oligonucleotides.

It will be understood by a skilled person that numerous different polynucleotides and nucleic acids can encode the same polypeptide as a result of the degeneracy of the genetic code. In addition, it is to be understood that skilled persons may, using routine techniques, make nucleotide substitutions that do not affect the polypeptide sequence encoded by the polynucleotides described here to reflect the codon usage of any particular host organism in which the polypeptides are to be expressed.

The polynucleotides described herein may comprise DNA or RNA. They may be single-stranded or double-stranded. They may also be polynucleotides that include within them synthetic or modified nucleotides. A number of different types of modification to oligonucleotides are known in the art. These include methylphosphonate and phosphorothioate backbones, addition of acridine or polylysine chains at the 3' and/or 5' ends of the molecule. For the purposes of the present invention, it is to be understood that the polynucleotides described herein may be modified by any method available in the art. Such modifications may be carried out in order to enhance the in vivo activity or life span of polynucleotides.

Where the polynucleotide is double-stranded, both strands of the duplex, either individually or in combination, are encompassed by the methods and compositions described herein. Where the polynucleotide is single-stranded, it is to be understood that the complementary sequence of that polynucleotide is also included.

The terms "variant," "homologue," or "derivative" in relation to a nucleotide sequence include any substitution of, variation of, modification of, replacement of, deletion of, or addition of one (or more) nucleotides from or to the sequence.

As indicated above, with respect to sequence identity, a "homologue" has at least 5% identity, at least 10% identity, at least 15% identity, at least 20% identity, at least 25% identity, at least 30% identity, at least 35% identity, at least 40% identity, at least 45% identity, at least 50% identity, at least 55% identity, at least 60% identity, at least 65% identity, at least 70% identity, at least 75% identity, at least 80% identity, at least 85% identity, at least 90% identity, or at least 95% identity to the relevant sequence shown in the sequence listings.

In particular, there is at least 95% identity, more particularly at least 96% identity, more particularly at least 97% identity, more particularly at least 98% identity, more particularly at least 99% identity. Nucleotide homology comparisons may be conducted using techniques well known in the art. One suitable sequence comparison program is the GCG Wisconsin Bestfit program, which is known in the art. The default scoring matrix has a match value of 10 for each identical nucleotide and -9 for each mismatch. The default gap creation penalty is -50 and the default gap extension penalty is -3 for each nucleotide. Various other methods are known in the art on how to calculate sequence identity.

In one embodiment, the estrogen-responsive 5'-regulatory nucleotide of the present invention has at least 98.1% or more sequence identity to a sequence corresponding to SEQ ID NO:2, SEQ ID NO:3, or SEQ ID NO:4. In particular, the estrogen-responsive 5'-regulatory nucleotide has 98.2% or more, 98.3% or more, 98.4% or more, 98.5% or more, 98.6% or more, 98.7% or more, 98.8% or more, 98.9% or more, 99.0% or more or 99.1% or more, 99.2% or more, 99.3% or more, 99.4% or more, 99.5% or more, 99.6% or more, 99.7% or more, 99.8% or more, 99.9% or more sequence identity to a sequence corresponding to SEQ ID NO:2, SEQ ID NO:3, or SEQ ID NO:4.

Polymerase chain reaction (PCR) may be employed during the processes involved in producing and detecting the transgenic fish of the present invention. Known methods of PCR include, but are not limited to, methods using paired primers, nested primers, single specific primers, degenerate primers, gene-specific primers, vector-specific primers, partially mismatched primers, and the like.

In one aspect, the present invention also relates to an expression cassette for use in producing the transgenic fish of the present invention. The expression cassette of the present invention includes the following: (i) a reporter nucleotide sequence encoding a reporter protein; (ii) an estrogen-responsive 5'-regulatory nucleotide sequence isolated from a medaka fish, wherein the regulatory nucleotide sequence is operably linked 5' to the reporter nucleotide sequence to induce expression of the reporter protein in the presence of estrogen or an estrogen-like compound; and (iii) a 3'-regulatory region operably linked 3' to the reporter nucleotide sequence.

In one embodiment, the expression cassette of the present invention includes a 5'-regulatory nucleotide sequence isolated from a brackish medaka fish of the species *Oryzias melastigma*. For example, the 5'-regulatory nucleotide sequence can be derived from the promoter region of a choriogenin H gene of *Oryzias melastigma*, where the choriogenin H gene has a nucleotide sequence of SEQ ID NO:1.

In one embodiment, the expression cassette of the present invention can include a 5'-regulatory nucleotide sequence that is derived from the promoter region of the *Oryzias melastigma* choriogenin H gene, where the nucleotide sequence corresponds to (i) SEQ ID NO:2 (corresponding to a -4827 bp omChgH5'-upstream region); (ii) SEQ ID NO:3

(corresponding to a -750 bp omChgH5'-upstream region); and (iii) SEQ ID NO:4 (corresponding to a -350 bp omChgH5'-upstream region).

In another embodiment, the reporter nucleotide sequence of the expression cassette of the present invention encodes a reporter protein that is an autofluorescent protein. Suitable examples of autofluorescent proteins can include, without limitation, an enhanced green fluorescent protein (EGFP), a green fluorescent protein (GFP), a blue fluorescent protein (BFP), a yellow fluorescent protein (YFP), a red fluorescent protein (RFP), and the like.

In one embodiment, the expression cassette of the present invention can include a 3'-regulatory region that comprises an mRNA polyadenylation signal. A suitable mRNA polyadenylation signal can include, without limitation, the 3'-flanking region of a choriogenin H gene of a medaka fish (e.g., *Oryzias melastigma* and *Oryzias latipes*). In particular, the mRNA polyadenylation signal can include, without limitation, the 3'-flanking region of the choriogenin H gene that includes the 3'-UTR (untranslated region) and the polyadenylation tail sequences of the choriogenin H gene. More particularly, the 3'-regulatory region can include the nucleotide sequence of SEQ ID NO:5.

The present invention also relates to a transformation vector that contains the expression cassette of the present invention.

The present invention also relates to a host cell transduced with the expression cassette of the present invention.

The present invention also relates to a transgenic cell of an aquatic animal, where the transgenic cell includes at least one expression cassette of the present invention. As set forth herein, the transgenic cell can be from an aquatic animal that is a freshwater, saltwater, and brackish water fish. Suitable examples of such fish are discussed herein and include, for example, *Oryzias* species and *Danio* species.

The present invention also relates to a transgenic fish that contains at least one expression cassette of the present invention. The transgenic fish of the present invention exhibits an observable mark when exposed to a liquid medium containing an estrogen or an estrogen-like compound. The observable mark includes the reporter protein expressed by the at least one expression cassette. The observable mark is visible in vivo or in vitro. The transgenic fish can be of a freshwater, saltwater, or brackish water fish. As noted, suitable examples of such fish are discussed herein and include, for example, *Oryzias* species and *Danio* species. In a particular embodiment, the transgenic fish is at a stage of development that includes, without limitation, an embryo stage, a larva stage, and/or an adult stage. More particularly, the transgenic fish can be a male adult.

The "expression cassette" of the present invention can also be referred to as a "recombinant construct." Recombinant constructs are generally known in the art to contain an expression control sequence (e.g., the estrogen-responsive 5'-regulatory nucleotide sequence), which is operably linked to a reporter sequence. That means that a polynucleotide comprising an expression control sequence of interest is cloned in a recombinant construct, such that the expression control sequence is operably linked to a reporter sequence. In particular, the reporter is a heterologous sequence.

The methods for making the recombinant constructs are conventional. Such methods, as well as many of the other molecular biological methods used in conjunction with the present invention, are discussed, e.g., in Sambrook, et al. (1989), *Molecular Cloning, a Laboratory Manual*, Cold Harbor Laboratory Press, Cold Spring Harbor, N.Y.; Ausubel et al. (1995). *Current Protocols in Molecular Biology*, N.Y.,

John Wiley & Sons Davis et al. (1986), Basic Methods in Molecular Biology, Elsevier Sciences Publishing, Inc., New York; Hames et al. (1985), Nucleic Acid Hybridization, IL Press; Dracopoli et al. Current Protocols in Human Genetics, John Wiley & Sons, Inc.; and Coligan et al. Current Protocols in Protein Science, John Wiley & Sons, Inc.

Suitable reporter sequences will be evident to those of skill in the art. For example, a suitable reporter protein can include autofluorescent proteins and enzymes detectable by a histochemical method.

As indicated elsewhere herein, the fluorescent protein can include, without limitation, a green fluorescent protein (GFP), an enhanced green fluorescent protein (EGFP), a red fluorescent protein (CFP and Red FP, RFP), a blue fluorescent protein (BFP), a yellow fluorescent protein (YFP), and fluorescent variants of these proteins. The heterologous fluorescent gene may be, for example, a gene encoding DsRed2, ZsGreen1, and ZsYellow1. The heterologous fluorescent gene may also be any variation or mutation of these genes, encoding fluorescent proteins including green fluorescent protein (GFP), enhanced green fluorescent protein (eGFP), yellow fluorescent protein (YFP), enhanced yellow fluorescent protein (eYFP), blue fluorescent protein (BFP), enhanced blue fluorescent protein (eBFP), cyan fluorescent protein (CFP), and enhanced cyan fluorescent protein (eCFP).

According to another embodiment, the enzyme that is detectable by a histochemical method is chosen from the group composed of luciferase,  $\beta$ -galactosidase,  $\beta$ -glucuronidase, alkaline phosphatase, chloramphenicol acetyl transferase, and alcohol dehydrogenase. According to a particular embodiment, it is luciferase. The term "luciferase" is intended to denote all the proteins which catalyze or initiate a bioluminescent reaction in the presence of a substrate called luciferin. The luciferase according to the invention may come from many organisms or systems that generate bioluminescence (see U.S. Pat. No. 6,152,358). For example, the luciferase according to the invention may come from *Renilla* (U.S. Pat. No. 5,418,155 and U.S. Pat. No. 5,292,658) or from *Photinus pyralis* or from *Luciola cruciata* (U.S. Pat. No. 4,968,613).

Techniques to detect protein reporters, either directly (e.g., by measuring the amount of reporter mRNA) or indirectly (e.g. by measuring the amount and/or activity of the reporter protein) are conventional. Many of these methodologies and analytical techniques can be found in such references as Current Protocols in Molecular Biology, F. M. Ausubel et al., eds., (a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc.), Enzyme Immunoassay, Maggio, ed. (CRC Press, Boca Raton, 1980); Laboratory Techniques in Biochemistry and Molecular Biology, T. S. Work and E. Work, eds. (Elsevier Science Publishers B. V., Amsterdam, 1985); Principles and Practice of Immunoassays, Price and Newman, eds. (Stockton Press, NY, 1991); and the like.

For example, changes in nucleic acid expression can be determined by polymerase chain reaction (PCR), ligase chain reaction (LCR), Q $\beta$ -replicase amplification, nucleic acid sequence based amplification (NASBA), and other transcription-mediated amplification techniques; differential display protocols; analysis of northern blots, enzyme linked assays, micro-arrays and the like. Examples of these techniques can be found in, for example, PCR Protocols A Guide to Methods and Applications (Innis et al., eds, Academic Press Inc. San Diego, Calif. (1990)).

In a particular embodiment, the amount and/or activity of a reporter expression product (e.g., a protein) is measured. A

fluorescent marker, such as EGFP, can be detected by detecting its fluorescence in the cell (e.g., in a brackish medaka fish embryo). For example, fluorescence can be observed under a fluorescence microscope. Reporters such as EGFP, which are directly detectable without requiring the addition of exogenous factors, are preferred for detecting or assessing gene expression during fish embryonic development. A transgenic fish embryo carrying a recombinant construct of the invention encoding an EGFP reporter can provide a rapid real time in vivo system for analyzing spatial and temporal expression patterns of developmentally regulated liver genes, and for determining the presence of an estrogenic compound.

The recombinant construct of the present invention can be cloned into a suitable vector. The vector can then be used, e.g., to propagate the recombinant construct. Generally, before introducing a recombinant construct of the invention into a fish embryo, it is desirable to remove the vector sequences. Preferably, the vector/construct is designed so that the recombinant construct can be excised with one or two appropriate restriction enzymes.

Large numbers of suitable vectors are known to those of skill in the art, and many are commercially available. The following vectors are provided by way of example: (i) bacterial vectors, including pQE70, pQE60, pQE-9 (Qiagen), pBS, pD10, phagescript, psiX174, pBluescript SK, pBSKS, pNH8A, pNH16a, pNH18A, pNH46A (Stratagene), pTRC99a, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia); and (ii) eukaryotic vectors, including pWLNEO, pSV2CAT, pOG44, pXT1, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia). However, any other plasmid or vector may be used as long as it is replicable and viable in the host.

In one aspect, the present invention provides a method for detecting the promoter activity in a cell of an estrogen-responsive 5'-regulatory nucleotide sequence of the present invention. This can be done in a eukaryotic cell by introducing the recombinant construct into the eukaryotic cell, and detecting the presence and/or activity of the reporter sequence in the cell. A variety of eukaryotic cells can be used; suitable cells will be evident to the skilled worker. In particular embodiments, the reporter sequence encodes EGFP, and the eukaryotic cell is in or from a fish, such as a brackish medaka fish, or is in or from a brackish medaka fish embryo.

Many art-recognized methods are available for introducing polynucleotides, such as the constructs of the present invention, into cells. The conventional methods that can be employed, include, e.g., transfection (e.g., mediated by DEAE-Dextran or calcium phosphate precipitation), infection via a viral vector (e.g., retrovirus, adenovirus, adenovirus-associated virus, lentivirus, pseudotyped retrovirus or poxvirus vectors), injection, such as microinjection, electroporation, sonoporation, a gene gun, liposome delivery (e.g., Lipofectin®, Lipofectamine® (GIBCO-BRL, Inc., Gaithersburg, Md.), Superfect® (Qiagen, Inc. Hilden, Germany) and Transfectam® (Promega Biotec, Inc., Madison, Wis.), or other liposomes developed according to procedures standard in the art), or receptor-mediated uptake and other endocytosis mechanisms.

Methods for introducing the recombinant construct into a fish embryo are discussed in more detail elsewhere herein.

As used herein, "transgenic fish" refers to fish, or progeny of a fish, into which an exogenous recombinant construct has been introduced. A fish into which a construct has been introduced includes fish that have developed from embryonic cells into which the construct has been introduced. As used herein, an exogenous construct is a nucleic acid that is artificially introduced, or was originally artificially introduced, into an

animal. The term artificial introduction is intended to exclude introduction of a construct through normal reproduction or genetic crosses. That is, the original introduction of a gene or trait into a line or strain of animal by cross breeding is intended to be excluded.

However, fish produced by transfer, through normal breeding, of an exogenous construct (that is, a construct that was originally artificially introduced) from a fish containing the construct are considered to contain an exogenous construct. Such fish are progeny of fish into which the exogenous construct has been introduced. As used herein, progeny of a fish are any fish which are descended from the fish by sexual reproduction or cloning, and from which genetic material has been inherited. In this context, cloning refers to production of a genetically identical fish from DNA, a cell, or cells of the fish. The fish from which another fish is descended is referred to as a progenitor fish. As used herein, development of a fish from a cell or cells (embryonic cells, for example), or development of a cell or cells into a fish, refers to the developmental process by which fertilized egg cells or embryonic cells (and their progeny) grow, divide, and differentiate to form an adult fish.

In one embodiment, a transgenic fish of the present invention is one whose somatic and germ cells contain at least one genomically integrated copy of a recombinant construct of the invention. The invention further provides a transgenic fish gamete, including a transgenic fish egg or sperm cell, a transgenic fish embryo, and any other type of transgenic fish cell or cluster of cells, whether haploid, diploid, triploid or other zygosity having at least one genomically integrated copy of a recombinant construct of the present invention.

As used herein, the term "embryo" includes a single cell fertilized egg (i.e., a zygote) stage of the organism. In particular, the recombinant construct is integrated into the fish's somatic and germ cells such that it is stable and inheritable (is stably transmitted through the germ line). The transgenic fish or fish cell preferably contains a multiplicity of genomically integrated copies of the construct; more specifically, the multiple copies of the construct are integrated into the host organism's genome in a contiguous, head-to-tail orientation.

Progeny of the transgenic fish containing at least one genomically integrated copy of the construct, and transgenic fish derived from a transgenic fish egg, sperm, embryo or other fish cell of the present invention, are also included in the present invention. A fish is "derived from" a transgenic fish egg, sperm cell, embryo or other cell if the transgenic fish egg, sperm cell, embryo or other cell contributes DNA to the fish's genomic DNA. For example, a transgenic embryo of the present invention can develop into a transgenic fish of the present invention; a transgenic egg of the present invention can be fertilized to create a transgenic embryo of the present invention that develops into a transgenic fish of the present invention; a transgenic sperm cell of the present invention can be used to fertilize an egg to create a transgenic embryo of the present invention that develops into a transgenic fish of the present invention; and a transgenic cell of the present invention can be used to clone a transgenic fish of the present invention. In some embodiments of the present invention, the transgenic fish is sterile. The present invention further includes a cell line derived from a transgenic fish embryo or other transgenic fish cell of the present invention, which contains at least one copy of a recombinant construct of the present invention. Methods of isolating such cells and propagating them are conventional.

In one embodiment, the disclosed transgenic fish of the present invention are produced by introducing a recombinant construct of the present invention into cells of a fish, particu-

larly into embryonic cells, and more particularly into a single cell embryo. Where the transgene construct is introduced into embryonic cells, the transgenic fish is obtained by allowing the embryo to develop into a fish. Introduction of constructs into embryonic cells of fish, and subsequent development of the fish, are simplified by the fact that embryos develop outside of the parent fish.

The disclosed recombinant constructs can be introduced into embryonic fish cells using any suitable technique. Many techniques for such introduction of exogenous genetic material have been demonstrated in fish and other animals. These include microinjection (described by, for example, Culp et al. (1991) *Proc Natl Acad Sci USA* 88, 7953-7957), electroporation (described by, for example, Inoue et al. (1990), *Cell Differ. Develop.* 29, 123-128; Muller et al. (1993), *FEBS Lett.* 324, 27-32; Murakami et al. (1994), *J. Biotechnol.* 34, 35-42; Muller et al. (1992), *Mol. Mar. Biol. Biotechnol.* 1, 276-281; and Symonds et al. (1994), *Aquaculture* 119, 313-327), particle gun bombardment (Zelenin et al. (1991), *FEBS Lett.* 287, 118-120), retroviral vectors (Lu et al (1997). *Mol Mar Biol Biotechnol* 6, 289-95), and the use of liposomes (Szelei et al. (1994), *Transgenic Res.* 3, 116-119).

Embryos or embryonic cells can generally be obtained by collecting eggs immediately after they are laid. It is generally preferred that the eggs be fertilized prior to or at the time of collection. This is particularly accomplished by placing a male and female fish together in a tank that allows egg collection under conditions that stimulate mating. After collecting eggs, one technique is that the embryo be exposed for introduction of genetic material by removing the chorion. This can be done manually or, particularly, by using a protease such as pronase. A fertilized egg cell prior to the first cell division is considered a one-cell embryo, and the fertilized egg cell is thus considered an embryonic cell.

After introduction of the transgene construct, the embryo is allowed to develop into a fish. This generally need involve no more than incubating the embryos under the same conditions used for incubation of eggs. However, the embryonic cells can also be incubated briefly in an isotonic buffer. If appropriate, expression of an introduced transgene construct can be observed during development of the embryo.

Fish harboring a transgene can be identified by any suitable means. For example, the genome of potential transgenic fish can be probed for the presence of construct sequences. To identify transgenic fish actually expressing the transgene, the presence of an expression product can be assayed. Several techniques for such identification are known and used for transgenic animals and most can be applied to transgenic fish. Probing of potential or actual transgenic fish for nucleic acid sequences present in or characteristic of a transgene construct can be accomplished by Southern or Northern blotting. Also, detection can be achieved using polymerase chain reaction (PCR) or other sequence-specific nucleic acid amplification techniques.

After "founder" transgenic fish are identified, one can mate them to wild type fish to identify those fish which comprise the transgene in their germ cells. Transgenic fish of the present invention can be either male or female. A transgenic fish of the present invention can be hemizygous for the transgene, which is a particular state for maintenance of transgenic fish lines. Alternatively, hemizygous fish can be crossed with each other to produce homozygous fish or fish lines. Homozygous diploids can also be produced by other methods, e.g., interruption of the second meiotic divisions with hydrostatic pressure using a French press.

The disclosed recombinant constructs can be integrated into the genome of the fish. However, the disclosed transgene

construct can also be constructed as an artificial chromosome. Such artificial chromosomes containing more than 200 kb have been used in several organisms. Artificial chromosomes can be used to introduce very large transgene constructs into fish. This technology is useful since it can allow faithful recapitulation of the expression pattern of genes that have regulatory elements that lie many kilobases from coding sequences.

In another embodiment, the present invention includes a genomically identical population of transgenic fish, each of whose somatic and germ cells contain at least one genomically integrated copy of a recombinant construct of the present invention. The genomically identical population is a unisex population and can be male or female. Particular embodiments of the genomically identical transgenic fish population are essentially as described for the transgenic fish of the present invention. In an alternative embodiment, the present invention includes a population of transgenic fish, i.e., an in-bred line, the members of which are not necessarily genomically identical but are homozygous with respect to genomically integrated constructs.

The present invention also relates to various methods of using the transgenic fish of the present invention. In particular embodiments, these methods involve the detection of the presence or absence of an estrogenic compound in a medium, particularly in a liquid medium. More particularly, such methods are based on the induced expression of the reporter gene (also referred to herein as the reporter nucleotide sequence) that is operably linked to an estrogen-responsive 5'-regulatory nucleotide sequence of the present invention (as described elsewhere herein). Generally, when a transgenic fish of the present invention is exposed to an estrogenic compound, the estrogen-responsive 5'-regulatory nucleotide sequence will induce the expression of the reporter gene. Expression of the reporter gene results in an observable mark on the transgenic fish, indicating the presence of an estrogenic compound. As used herein, an "estrogenic compound or substance" refers to estrogen or any estrogen-like compound or substance.

As used herein, a "liquid medium" can refer to a number of different liquids. For example, it can include a liquid or water sample into which a test substance of interest has been added. It can also include a liquid or water sample into which a test substance of interest has not been added. Particular examples of a liquid medium suitable for use in the methods of the present invention can include freshwater, brackish water, and saltwater from any liquid or water source (e.g., rivers, lakes, bays, ponds, streams, oceans, etc.), whether these sources are natural or man-made.

Some of the various methods of using the transgenic fish of the present invention are generally provided below and more specifically provided in the Examples section hereof.

One method of the present invention relates to a method of screening a liquid medium for an estrogenic substance. This method involves exposing a transgenic fish of the present invention to a liquid medium to be tested (i.e., to be tested for the presence of an estrogenic substance). After this exposing step, the method then involves determining whether or not the transgenic fish exhibits an observable mark produced by the induced expression of the reporter gene (contained in the transgenic fish). The presence of the observable mark indicates that the liquid medium contains an estrogenic substance.

In one embodiment of this method, the determining step can include in vivo autofluorescence microscopy, where the reporter gene encodes an autofluorescent reporter protein.

In another embodiment of this method, the method can further include quantifying the estrogen-like activity of a test substance identified to be an estrogenic substance according to the determining step, where the test substance is identified as being an estrogenic substance if the transgenic fish exhibits an observable mark.

In another embodiment of this method, the quantifying step includes (i) generating at least one image of the observable mark; and (ii) subjecting the at least one image of the observable mark to image analysis software to quantify the signal intensity of the observable mark.

Another method of the present invention relates to a method of screening for a compound having anti-estrogenic activity. This method involves the following steps: (a) providing a first transgenic fish and a second transgenic fish of the present invention, where the first and second transgenic fish are of the same species and at substantially the same developmental stage; (b) exposing the first transgenic fish to a first liquid medium, where the first liquid medium includes an estrogen or estrogen-like compound; (c) exposing the second transgenic fish to a second liquid medium, where the second liquid medium includes the first liquid medium and a test compound; and (d) comparing the quantified intensity of any observable mark exhibited by the first transgenic fish with the quantified intensity of any observable mark exhibited by the second transgenic fish, where a decrease in the quantified intensity of the observable mark in the second transgenic fish compared to that of the first transgenic fish indicates that the test compound has anti-estrogenic activity.

Another method of the present invention relates to a method for investigating the effect of an estrogenic compound on liver regeneration. Generally, this method involves performing a partial hepatectomy on an adult transgenic fish of the present invention, where the partial hepatectomy is effective to remove a portion of the liver of the transgenic fish; exposing the transgenic fish to a test liquid medium containing a test estrogenic compound; and analyzing the liver of the transgenic fish to detect any regeneration of liver tissue.

In one embodiment of this method, the analyzing step can include comparing liver regeneration parameters of the transgenic fish taken at the following stages: (i) before the partial hepatectomy; (ii) after the partial hepatectomy but before the exposing step; and (iii) after the partial hepatectomy and after the exposing step. Suitable liver regeneration parameters can include, without limitation, liver volume, liver shape, liver weight, and/or liver-to-body weight ratio. In one aspect, determining the presence and/or intensity of the observable mark exhibited by the transgenic fish is effective to assist in measuring the liver regeneration parameters. In a particular embodiment, the liver shape and liver volume parameters are measured by generating three-dimensional images of the liver using confocal microscopy combined with image analysis software.

Another method of the present invention relates to a method for investigating the effect of different estrogenic compounds on liver regeneration, as opposed to investigating a single estrogenic compound. Generally, this method involves the following steps: (a) providing a first transgenic fish and a second transgenic fish of the present invention, where the first and second transgenic fish are adults of the same species; (b) exposing the first transgenic fish to a first liquid medium that includes a first test estrogenic compound solution; (c) exposing the second transgenic fish to a second liquid medium comprising a second test estrogenic compound solution; (d) quantifying the intensity of any observable mark exhibited by the first transgenic fish and the second transgenic fish; (e) performing a partial hepatectomy on the

first and second transgenic fish, where the partial hepatectomy is effective to remove a portion of the liver of the first and second transgenic fish; (f) repeating steps (b) through (d) of this method; and (g) analyzing the liver of the first and second transgenic fish to compare the effects of the estrogenic compounds contained in the first liquid medium and the second liquid medium on any regeneration of liver tissue.

In one embodiment of this method, the analyzing step can include comparing liver regeneration parameters of the transgenic fish taken at the following stages: (i) before the partial hepatectomy; (ii) after the partial hepatectomy but before the exposing step; and (iii) after the partial hepatectomy and after the exposing step. Suitable liver regeneration parameters can include, without limitation, liver volume, liver shape, liver weight, and/or liver-to-body weight ratio. In one aspect, determining the presence and/or intensity of the observable mark exhibited by the transgenic fish is effective to assist in measuring the liver regeneration parameters. In a particular embodiment, the liver shape and liver volume parameters are measured by generating three-dimensional images of the liver using confocal microscopy combined with image analysis software.

In another embodiment of this method, the method can further include separating hepatocytes exhibiting observable marks from other liver cells not exhibiting observable marks, where the separating is conducted using flow cytometry.

In another embodiment of this method, the method can further include conducting hepatocyte metabolomic and proteomic analyses of the liver of the first and second transgenic fish.

Provided below are particular embodiments of various aspects of the present invention, some of which have been generally described above.

#### Cloning of Brackish Medaka-Derived Choriogenin H Gene

In a particular embodiment of the present invention, the brackish medaka fish to be used in cloning a choriogenin H gene (omChgH) and the medaka fish to be used in introduction of the constructed transgenic gene are not particularly limited, as long as they belong to a species *Oryzias melastigma* (alternative name *Oryzias dancena*). The medaka fish used in this embodiment of the present invention were obtained from a commercial hatchery in Taiwan. The obtained medaka fish can be maintained in freshwater, brackish water, as well as seawater. In this embodiment of the present invention, the fish were maintained in 30 ppt artificial seawater at  $28 \pm 1^\circ \text{C}$ . with a constant light cycle of 14 h-light/8 h-dark and fed with commercial hormone-free flake food and brine shrimp (*Anemia salina*).

The *Oryzias melastigma*-derived estrogen-responsive choriogenin H gene promoter region used in this embodiment of the present invention is captured after sequential genome walks in Dra I and EcoR V genomic DNA libraries according to the manufacturer's instruction on the genome walking kit (BD Biosciences).

In this embodiment of the present invention, choriogenin H 3'-flanking region includes choriogenin H 3'-untranslated region (represented as UTR) and partial genomic DNA sequence downstream is isolated. To clone 3'-UTR downstream genomic sequence, choriogenin H exon-intron structure is identified. The choriogenin H 3'-downstream genomic nucleotide is isolated using inverse-PCR (polymerase chain reaction). A sequence of ca. 800 bp choriogenin H 3'-flanking

region from transcription terminal is isolated and deposited at GenBank with Accession No. DQ778335.

#### Construction of Transgene Plasmid

In one embodiment of the present invention, the *Oryzias melastigma*-derived choriogenin H promoter region cloned by the aforementioned method, EGFP cDNA sequence (i.e., the reporter nucleotide sequence), and *Oryzias melastigma*-derived choriogenin H 3'-flanking region cloned by the aforementioned method, or poly (A) signal of other genes, are introduced into a vector. The introduction of these nucleotide sequences into a vector can be performed in accordance with a known genetic engineering process. In this manner, it is possible to construct a recombinant vector into which the *Oryzias melastigma*-derived choriogenin H promoter region linked with EGFP encoding region are inserted.

In this and other embodiments of the present invention, the vector to be used is not particularly limited, as long as the promoter sequence, encoding sequence, and poly (A) signal sequence are operably inserted.

#### Production of Transgenic Brackish Medaka Fish

In one embodiment of the present invention, the recombinant transgene plasmid constructed as mentioned above is linearized and transferred into the plasma of brackish medaka fertilized eggs, thereby making the transgenic brackish medaka fish capable of expressing the EGFP regulated by the inserted omChgH promoter in an estrogen-dependent manner, just like its internal omChgH does. As the brackish medaka fertilized egg to be transferred in this and other embodiments of the present invention, embryos at one-cell stage within half an hour after fertilization are used. The recombinant transgene sequence can be transferred into the single-cell of fertilized eggs by means of microinjection.

To get one-cell stage brackish medaka eggs, a mating tank system can be used to separate male and female fish the day before microinjection. Male and female fish are released to mate by pulling the separator away from the mating tank before proceeding microinjection. In an embodiment of the present invention, the fertilized eggs which have been subjected to microinjection are raised at  $28 \pm 1^\circ \text{C}$ . for 2-3 months till they become adult fish.

Adult fish having the introduced gene integrated into its genome can be screened by the following methods. First, adult fish mate with wild fish and DNA is extracted from their eggs. The extracted DNA is then PCR amplified using EGFP specific primers, and the amplified product is subjected to electrophoresis. Second, adult fish mate with wild fish. Their eggs are raised to hatch out and then exposed to estrogen or estrogen-like chemical substances overnight or longer. These larvae are then checked under fluorescent microscope equipped with GFP filter set to see whether there is GFP expression in the liver. By these two methods, brackish medaka fish having the introduced nucleotide sequence integrated into the genome of a germ cell (sperm or egg) can be successfully identified.

Subsequently, the brackish medaka fish containing the introduced nucleotide sequence into the genome of a germ cell is mated with wild type fish. Their progeny inherit the introduced nucleotide sequence and can be screened for the introduced nucleotide sequence by the aforementioned two methods. This fish can pass the transgene from generation to generation. Such fish include a desired transgenic brackish medaka fish of the present invention.

Without any treatment, the expression of the introduced EGFP in the transgenic brackish medaka is restricted to the liver of reproducing females that synthesize high concentration of estrogen, and no EGFP expression is observed in the liver of embryos, larvae and male fish. However, when the transgenic brackish medaka fish is exposed to estrogen or estrogen-like chemical substances, the EGFP transgene can also be induced to express in the liver of embryo, larvae and male fish. The expression profile of the introduced EGFP is consistent with the internal expression profile of omChgH in this brackish medaka (X. Chen et al. *Ectotoxicol. Environ. Saf.* 71:200-208 (2008)), except that omChgH protein is a secretory protein that will transport to ovary through blood circulation after its synthesis in the liver, while EGFP is not a secretory protein and will stay in the liver after its synthesis. As the transgenic brackish medaka female fish express EGFP without any outer induction, non-EGFP-expressing embryo, larvae and male transgenic brackish medaka fish are preferably to be used to test estrogen like activity.

Due to high translucency of the brackish medaka fish during embryo and early larvae stage, the response of the transgenic brackish medaka fish to estrogen or estrogen-like substances at these stages can be easily observed in vivo using a fluorescence microscope. The early stage larvae (before 15 days past fertilization), which are most translucent, are a particularly good stage for testing estrogen-like activity in vivo. Further, if the estrogen or estrogen-like chemicals treated transgenic brackish medaka fish are transferred to clean water, the induced EGFP signal will gradually disappear in around one week, and the EGFP can be re-induced if the fish is put back to solution containing estrogen or estrogen-like chemical substances.

#### Method of Detecting Estrogen-Like Activity

In one embodiment of the present invention, whether estrogen or estrogen-like substances are present or not in a test water sample can be easily detected by exposing the transgenic brackish medaka fish of the present invention into the test water for a period of time and then observing the exposed transgenic fish under fluorescence microscope to see whether EGFP expression is induced in the liver or not.

As used herein, "test water" can be, but is not restricted to, water collected from the environment (e.g., river, estuary, brackish, etc.), which may contain one or more than one estrogen-like substance, or water added with one substance (e.g., pharmaceuticals, cosmetics, food stuff, etc.), which may contain estrogen or estrogen-like substances, or water added with one or more than one chemical to be tested, which may act separately, or in combination, like estrogen.

When testing a potential estrogenic chemical or mixture substance, in one embodiment of the present invention, testing can be done in a serial of concentrations, since each chemical substance has varying toxicity as well as potential estrogen-like activity; additionally, it is also suitable and useful to include an estrogen (E2) exposure of known concentration as a reference to measure the estrogen-like activity and calculate the estrogen equivalency of the test chemical substance. Exposing the transgenic brackish medaka fish to the test water for 1-2 days will be long enough for most samples. The exposure condition may be the same as that of raising brackish medaka fish.

One advantage of this method is that estrogen-like activity of test water can be quantified by measuring the induced EGFP signal intensity in the liver. After appropriate time of exposure, fish exposed to test water of different estrogen-like activity will express varying quantities of EGFP in the liver,

which can be easily differentiated by observation under fluorescence microscope. In one embodiment, to quantify the induced EGFP signal, fluorescence image of the EGFP expressing liver will be recorded using a camera connected to the fluorescence microscope, and the EGFP signal intensity in the liver area can be analyzed by image analysis software. When exposure to estrogen of known concentration is also included as reference, the estrogen-like activity of the test water can be converted to estrogen potency equivalent.

As embodiments of the transgenic brackish medaka fish of the present invention can be used to easily monitor the estrogen-like activity of test water, this method can be used to rapidly screen potential estrogenic chemical substances, as well as evaluate the combined estrogen-like activity of chemical mixtures. When chemical substances are tested separately as well as in combination, the mode of action of the chemical substances can be predicted. For example, some chemical substances may act estrogenically and some may act anti-estrogenically. When chemical substances of the same action mode, for example, estrogenic mode, are tested together, their total estrogen-like activity will be higher than that of any one of them alone. However, when estrogenic and anti-estrogenic chemical substances are mixed together, the final estrogen-like activity of the mixture will be lower than that of the estrogenic chemical substance.

#### Advantages of the Transgenic Brackish Medaka Fish Detection System

The various aspects of the present invention provide a number of advantages. Some of these advantages are described below, and such descriptions are not meant to include all of the advantages.

First, there are many advantages for using the transgenic fish of the present invention to detect estrogenic agents or events. The aqueous environment is the ultimate sink for natural and anthropogenic chemicals (J. P. Sumpter, *Toxicol. Lett.* 102-103:337-342 (1998)). Fish inhabit in water are environmentally relevant models for health risk assessment of aquatic environments. Despite the evolutionary distance between fish and humans, fish share most developmental pathways, physiological mechanisms and organ systems with humans (A. R. Cossins and D. L. Crawford, *Nature Rev. Genet.* 6:324-333 (2005)); and, thus, occupy a prominent position in the field of toxicology, development as well as biomedical studies. Fish are easy to handle, manipulate and observe, and are increasingly being used as an alternative, non-mammalian animal model to reduce or replace traditional mammalian models in research and testing. Fish, particularly small fish such as Japanese medaka (*Oryzias latipes*) and zebrafish (*Danio rerio*), have been widely used as laboratory model species in different fields of research. Advances made with these two aquarium fish models have led to the complete sequencing of their genomes (E. Ekker et al., *Zebrafish* 4, 239-251 (2003); M. Kasahara et al., *Nature*, 447:714-719 (2007)), and numerous transgenic fish strains have been established for these two research models, several of which have been aimed to detect estrogen-like activity as the present invention does (T. Kawamura, *Zoolog. Sci.* 19(2): 1355-1361 (2002); T. Ueno et al., *Mech. Dev.* 121(7-8):803-15 (2004); K. Krauchi et al., *Environ. Sci. Technol.* 39(8): 2762-8 (2005); Z. Zeng et al., *Environ. Sci. Technol.* 39(22): 9001-8 (2005); T. Hano et al., *Environ. Sci. Technol.* 41(4): 1473-9 (2007); M. A. Salam et al., *J Environ Sci Health A Tox Hazard Subst Environ Eng.* 43(3):272-7 (2008); J. Legler et al., *Environ. Sci. Technol.* 34:4439-4444 (2000); S. K. Tong et al., *Genesis*, 47(2):67-73 (2009)).



Second, it is advantageous to use fluorescence protein gene as reporter gene, through which the estrogen-like activity can be monitored by in vivo measurement of reporter gene expression without sacrificing the fish. Of these transgenic strains developed to detect estrogen-like activity, 3 strains used fluorescence protein reporter gene (T. Ueno et al., *Mech. Dev.* 121(7-8):803-15 (2004); K. Krauchi et al., *Environ. Sci. Technol.* 39(8):2762-8 (2005); Z. Zeng et al., *Environ. Sci. Technol.* 39(22):9001-8 (2005)), and have demonstrated its incomparable advantage for in vivo monitoring. However, none of these strains is sensitive enough to monitor the estrogen-like activity of environmental water samples without any concentration processing.

Third, the host organism can be selected to provide various advantages. For example, brackish medaka (*Oryzias melastigma*) shares most of the advantages that laboratory models zebrafish (*Danio rerio*) and Japanese medaka (*Oryzias latipes*) have, while possessing the advantage to thrive in varying salinity environment ranging from freshwater to seawater. The adaptability to a wide range of salinity makes this fish a universal model for both freshwater and seawater studies. And previous studies showed that research techniques and resources for this new model can be easily adapted from well established models such as zebrafish and Japanese medaka (X. Chen et al., *Comp. Biochem. Physiol. C. Toxicol. Pharmacol.* 149:647-655 (2009)).

Fourth, is that an estrogen-dependent liver-specific gene, omChgH, is identified and used in various embodiments of the present invention. omChgH is an egg envelope protein. The term "estrogen-dependent" used herein means this gene only expresses in the presence of estrogen or estrogen-like chemical substances. The term "liver-specific" used herein means the expression of this gene is restricted exclusively to the liver, not in any other organs. Under normal circumstances (in clean water without any treatment), this gene only expresses in reproducing female when internal estrogen concentration is high, but not in embryo, larvae and male fish (X. Chen et al. *Ectotoxicol. Environ. Saf.* 71:200-208 (2008)), in which internal estrogen level is almost undetectable. However, in the presence of external estrogen or estrogen-like chemical substances, this gene can also be induced to express in non-expressing embryo, larvae and male fish. Thus, using fluorescence protein gene as reporter marker regulated by the promoter of omChgH, the induced expression of omChgH gene will be represented as fluorescence protein in the liver, which can be easily detected under fluorescence microscope.

As noted above, in one aspect, the present invention relates to the development of a transgenic fish containing a genomically integrated nucleotide sequence comprising a fluorescence protein gene as reporter marker regulated by an estrogen-responsive promoter.

### EXAMPLES

The following examples, while exemplary of the present invention, are not to be construed as specifically limiting the invention. Accordingly, variations and equivalents, now known or later developed, that would be within the purview of one skilled in the art are to be considered to fall within the scope of this invention.

#### Example 1

##### Development of a Transgenic Fish

##### Preparation of Brackish Medaka Genomic DNA

Genomic DNA (represented as gDNA) was extracted from an adult female with intestine dissected out. The sample was

pestled in liquid nitrogen to small particles and incubated with 10 ml DNA extraction solution (150 mM NaCl, 0.5% sodium dodecyl sulfate (SDS), 25 mM ethylenediaminetetraacetic acid (EDTA), 10 mM Tris-HCl, pH 8.0) and 1% proteinase K at 45° C. for more than 4 h. Same volume phenol was added to the solution and mixed by slow rotation for 0.5 h. After 2,000 rpm centrifugation for 10 min, aqueous phase containing gDNA was transferred to a new polytron tube (50 ml) and mixed with the same volume of phenol/chloroform on a rotor for 10 min. Solution was then centrifuged at 2,000 rpm for 10 min, and aqueous phase was transferred to a new tube. gDNA solution was further purified by same volume of chloroform followed by 5 min centrifugation at 2,000 rpm. Aqueous phase was transferred to a new tube and gDNA was precipitated out by absolute ethanol. gDNA was carefully picked out using transfer tips, dissolved in nuclease-free water and kept at -20° C.

##### Cloning Brackish Medaka-Derived omChgH Gene Promoter Region

The 5'-flanking region of choriogenin H gene was captured after two rounds of genome walking according to the manufacturer's instructions (BD Biosciences). Nested PCR amplifications were performed using two adaptor-specific primers (GW-AP1: 5'-GTAATACGACTCACTATAGGGC-3' (SEQ ID NO:6), outer and GW-AP2: 5'-ACTATAGGGCACGCGTGGT-3' (SEQ ID NO:7), inner; provided in the kit) in combination with two anti-sense gene specific primers (represented as GSPs). GSPs used in the first genome walking in Dra I gDNA library were outer: 5'-AACGTGT-TGAGGGTCCTGCGGCTTC-3' (SEQ ID NO:8) and inner: 5'-CTGTGGATAGTATGGAGGGTATGGAACC-3' (SEQ ID NO:9); and GSPs for the second genome walking in EcoRV gDNA library were designed based on the result of the first genome walking, there were outer: 5'-GTGTAATGGATGTGGACTTTTCTATAAGACAACC-3' (SEQ ID NO:10) and inner: 5'-CCCAAAATGCATGTGCAGCTGATGGC-3' (SEQ ID NO:11). A 50-μl PCR reaction contained 5 μl of 10xPCR buffer, 1 μl of dNTP (10 mM), 1 μl of GSP (10 μM), 1 μl of adaptor primer (10 μM), 5 μl of 1/50 diluted adaptor-ligated genomic library (for primary PCR) or 1/500 diluted primary PCR product (for secondary PCR) and 1 μl of 50xBD Advantage 2 polymerase mix (BD Biosciences). The amplification profile for both primary and secondary PCRs consisted of 5 cycles of 94° C. for 15 s and 72° C. for 3 min, 5 cycles of 94° C. for 15 s and 70° C. for 3 min, and 25 cycles of 94° C. for 15 s and 68° C. for 3 min. The second PCR products were separated by electrophoresis, and the sharpest band was isolated, purified using Wizard® SV Gel and PCR Clean-up System (Promega), cloned to pGEM-T Easy Vector (Promega) and identified by commercial sequencing service (Tech Dragon).

##### Cloning of Brackish Medaka-Derived omChgH Gene 3'-Flanking Region

Inverse-PCR was performed to clone omChgH 3'-flanking region. After restriction endonuclease analysis of cloned coding region genomic DNA sequence (GenBank Accession No. EF392365) of omChgH gene, Nde I gDNA library was constructed by Nde I (Takara) digestion followed by T4 DNA ligase (Takara) ligation. Forward (5'-GAAAATCT-GTTTCATCATGGTTCAG-3') (SEQ ID NO:12) and Reverse (5'-TTCACAACCTGGTACCCTGTCCTGG-3') (SEQ ID NO:13) primers were designed near omChgH coding region 3'-end and upstream 5'-Nde I cutting site, respec-

tively. Reactions were performed in a 50- $\mu$ L volume containing 5  $\mu$ L 10 $\times$ PCR buffer, 1  $\mu$ L dNTP (10  $\mu$ M), 1  $\mu$ L each primer (10  $\mu$ M) and 1  $\mu$ L Taq enzyme (5 U/ $\mu$ L; Takara). The PCR thermal profile consisted of an initial denaturation at 95° C. for 4 min, followed by 35 cycles of 95° C. for 30 s, 60° C. for 30 s and 72° C. for 1 min. The amplicon was identified using the same method as omChgH 5'-flanking region.

#### Preparation of Transgenic Plasmids

omChgH5'-EGFP-olChgH3' plasmids construction: Totally 6595 bp of choriogenin H 5'-upstream region (GenBank Accession No. EF392365) from transcription initiation site was captured after sequential genome walks in Dra I and EcoR V gDNA libraries. In this embodiment, the plasmid ChgH-GFP (K. Krauchi et al., Environ. Sci. Technol. 39(8): 2762-8 (2005)) was used as vector. In this vector, Japanese medaka (*Oryzias latipes*) derived choriogenin H promoter region (represented as olChgH5') and 3'-flanking region (represented as olChgH3') were integrated, between which EGFP coding region was ligated. For easier following, this vector is represented as olChgH5'-EGFP-olChgH3'. After restriction sites analysis of both vector and omChgH 5'-upstream region, omChgH 5'-upstream region from -4827 to +15 from transcription initiation site was amplified using primers with built-in restriction sites: Forward (5'-TGCATG GCATGC TTAATTAA CTGCAG CCCGGG GTCGAC TCGTAC-CTCCAAAACCCAAC-3') (SEQ ID NO:14), Sph I, Pac I, Pst I, Sma I and Sal I restriction sites were underlined sequentially) and reverse (5'-CTCCAGTGCCTTG CCATGGT-3') (SEQ ID NO:15); Nco I site was underlined and translation start code was in bold). PCR fragment was subcloned into Sph I and Nco I restriction enzyme cutting sites of the olChgH5'-EGFP-olChgH3' vector to construct omChgH5'-EGFP-olChgH3' plasmid.

Additionally, two deletion mutants containing omChgH 5'-upstream region from -750 to +15 and from -350 to +15 were also constructed by PCR amplification and enzyme digestion, respectively, using omChgH5'-EGFP-olChgH3' plasmid as a template. The primers used for PCR amplification were 5'-TCGA CTGCAGTGCTCTCACTCTATGGGGTTC-3' (SEQ ID NO:16) (Pst I site was underlined, forward) and 5'-CTC-CAGTGCCTTGCCATGGT-3' (SEQ ID NO:17) (Nco I site was underlined, reverse). The amplicon then replaced the -4827 omChgH5' region of omChgH5'-EGFP-olChgH3' at Pst I and Nco I sites of omChgH5'-EGFP-olChgH3'. -350 omChgH 5'-EGFP-olChgH3' fragment was obtained by Xho I and EcoR I digestion of -4827 omChgH 5'-EGFP-olChgH3' plasmid.

omChgH5'-EGFP-SV40 Plasmid Construction: SV40 polyadenylation signal (represented as SV40 polyA) was amplified from DsRed2-1 vector (Clontech) by 5'-AGCTAC-TAGTCCATCTACATGGCCAAGAAG-3' (SEQ ID NO:18) (forward) and 5'-ATTTTGCCGATTTCGGCCTATTGGT-3' (SEQ ID NO:19) (reverse) primers, and ligated to pGEM®-T Easy vector for amplification. After digestion of by Not I and EcoR I, SV40 polyA replaced olChgH3' region in omChgH5'-EGFP-olChgH3' plasmid to construct plasmid omChgH5'-EGFP-SV40.

omChgH5'-EGFP-omChgH3' plasmids construction: A sequence of ca. 800 bp omChgH 3'-flanking region from transcription terminal (GenBank Accession No. DQ778335) was cloned by inverse-PCR described previously. For omChgH5'-GFP-omChgH3' construction, omChgH 3'-UTR was amplified using forward (5'-TGCAGCGGCCGCTGT-GAACCGACAGAAG-3') (SEQ ID NO:20) with reverse (5'-

CACGGATGTGTGTGTTTACC-3') (SEQ ID NO:21) primers, and 3'-flanking region was cloned by Forward (5'-GAAAATCTGTTTCATCATGGTTCAG-3') (SEQ ID NO:22) with Reverse (5'-TTCACAACCTGGTACCCTGTC-CTGG-3') (SEQ ID NO:23) primers. 3'-UTR and 3'-flanking region were then ligated using BD In-Fusion™ Dry-Down PCR Cloning Kits (Clontech) according to the instruction. omChgH translation stop codon downstream region was then PCR amplified by forward (5'-TGCAGCGGCCGCTGT-GAACCGACAGAAG-3') (SEQ ID NO:24) in combine with Reverse (5'-TTCACAACCTGGTACCCTGTCCTGG-3') (SEQ ID NO:25) primers. Amplicon was then ligated into pGEM®-T Easy vector for amplification. This sequence was then subcloned into Not I and EcoR I sites of omChgH5'-GFP-omChgH3' or omChgH5'-GFP-SV40 plasmids to form omChgH5'-GFP-omChgH3' plasmids.

#### Microinjection of Fish Embryos

Fish embryos were microinjected generally in accordance with Kinoshita et al. (Aquaculture, 143(3-4): 267-276 (1996)) with modifications. In order to maximize incorporation frequency of the transgene and reduce the degree of mosaicism in the founders, transgene solution was microinjected into the plasma of fertilized eggs at the one-cell stage. For easy collection of newly fertilized eggs, a male and a female fish was separated in each mating tank by a separator the day before microinjection. Before microinjection, remove the separator and freshly fertilized eggs can be collected directly from the belly of female fish using a plastic dropper. Eggs were separately from each other by using two fine forceps grasping filaments of embryos and rotating against each other. Microinjection was performed as fast as possible and the timing was rigidly controlled within 30 min past fertilization.

Injections were performed with the aid of stereomicroscope or compound microscope, micromanipulator, and oil or gas pressurized injection apparatus. Embryos were held in interstice of self-made glass cement holder on a glass slide. Transgenic DNA constructs were linearized, dissolved in 10 mM phosphate buffer (pH 7.5) containing 1 mM EDTA, 137 mM NaCl, 3 mM KCl. Linearized transgenic DNA fragments were injected at concentration ranging from 10 to 50 ng/ $\mu$ L.

#### Promoter Activity of omChgH 5'-Upstream Region

To analyze a region responsible for omChgH gene regulation, fragments containing three different sizes ranging from 4827 bp to 350 bp of omChgH 5'-upstream region fused with EGFP reporter gene were injected into one-cell *O. melastigma* embryos. GFP transient expression was observed in most embryos initiated at 3 days after injection. Hatchlings survived from microinjected embryos were exposed to 5.1 nM E2 for 24 h, and GFP expression was observed using fluorescence stereomicroscope. The incidence of GFP-expressing hatchlings for each construct was calculated (FIG. 1). Though the percentage of GFP fluorescence-positive fish varied slightly, omChgH5'-EGFP-olChgH3' constructs containing 4827 bp (-4827) and 758 bp (-758) 5'-upstream region of omChgH gene, respectively, were suggested to have the same activity at promoting the estrogen-dependent liver-specific expression of the downstream EGFP gene. However, obvious decrease of promoter activity was observed from -758 to -350 omChgH5'-EGFP-olChgH3' constructs, suggesting the existence of certain cis-regulatory element(s) located from -758 to -350 of omChgH 5'-upstream region. While the presence of GFP-positive hatchlings demonstrated

that 350 bp omChgH 5'-upstream region had the ability to regulate the estrogen-dependent liver-specific expression of its downstream gene.

Computer analysis identified putative cis-elements including one 5'-ERE-half site (at nt. Position -18), one C/EBP binding site (at nt. Position -158) and one estrogen-response element (ERE; at nt. Position -253) located within the first 350 bp 5'-upstream region, and another C/EBP binding site (at nt. Position -494) between -758 and -350 region of omChgH gene, respectively. The promoter activity demonstrated by -350 omChgH5'-GFP-olChgH3' construct suggests that some or all the three putative cis-elements identified within the first 350 bp 5'-upstream region are important for the regulation of the estrogen-dependent liver-specific expression of the downstream gene. While the obvious decrease of GFP-positive hatchling incidence from -758 to -350 omChgH5'-GFP-olChgH3' construct indicates that the C/EBP binding site at nt. Position -494 of omChgH is important for high cis-regulation of the estrogen-dependent liver-specific expression of downstream gene.

#### Regulation Effects of omChgH 3'-Flanking Region

As to some genes (e.g. vasa, nano), the 3'-flanking region is indispensable for the stability and localization of their mRNAs (M. Tanaka et al., Proc. Natl. Acad. Sci. U.S.A., 98(5): 2544-2549 (2001); M. Kopranner et al., Genes Dev. 15(21):2877-2885 (2001); H. Kurokawa et al., Dev. Growth Differ. 48(3):209-221 (2006)). In this embodiment, the importance of homogenous 3'-flanking region in omChgH transgenic studies was analyzed. olChgH3' of omChgH5'-GFP-olChgH3' fragment was replaced by SV40 polyA and omChgH3'-flanking region, respectively. These fragments were then injected into one-cell *O. melastigma* embryos and observation of estrogen-dependent GFP expression during hatchling stage was carried out as previously described and the results are summarized in FIG. 2. Flanked by SV40 polyA, -4728 omChgH5'-GFP DNA fragment demonstrated the ability to regulate the estrogen-dependent liver-specific expression of GFP fluorescence, and the GFP-positive hatchling incidence was even a little higher than that when it was flanked by olChgH3' but lower than that when it was flanked by omChgH3'. Additionally, higher GFP-positive hatchling incidences also observed in -750 and -350 omChgH5'-GFP-olChgH3' constructs than that of their accordingly omChgH5'-GFP-olChgH3' constructs (FIG. 1). These strongly indicate that homogenous 3'-flanking region is important for the high promoter activity of omChgH5'-GFP transgenic constructs, and the putative elements identified within the 3'-flanking region may explain the difference.

Moreover, results showed that similar promoter activity was observed between -4728 and -758 omChgH5'-GFP-olChgH3' constructs, while obvious decrease of promoter activity was observed in -350 omChgH5'-GFP-olChgH3' construct (FIG. 1). This further confirmed previous results (FIG. 1) that 350 bp omChgH 5'-upstream region was sufficient for regulating the estrogen-dependent liver-specific expression of its downstream gene, while the C/EBP binding site at nt. Position -494 of omChgH enhances the promoter activity.

#### Identification of Transgenic Fish

Injected embryos were raised for ca. 2-3 months to adult fish. These founders (F0) were then mated with wild-type fish individually. Their embryos were collected for genomic DNA extraction using Accuprep genomic DNA Extraction Kit

(Bionner, Korea), and the presence of EGFP transgene was tested by PCR using EGFP specific primers (forward: 5'-TGCTGCCCCGACAACCACTACC-3' (SEQ ID NO:26) and reverse: 5'-TTACTTGTACAGCTCGTCCATGC-3' (SEQ ID NO:27)). More embryos of EGFP-positive founders (F0) were collected and progenies (F1) carrying the EGFP transgene were screened by green fluorescence mark after exposure to estrogen (17 $\beta$ -estradiol, 5.1 nM) during hatchling stage. F1 fish expressed EGFP in response to estrogen were stable germ-line transgenic fish that can transfer the transgene to their progenies. Totally 11 germ-line transgenic fish strains were identified. Exposed to the same concentration of estrogen, the transgenic fish strain that showed most sensitive to estrogen was selected for detail analysis.

#### Transgene EGFP Expression

Without any treatment, the expression of the introduced EGFP in the transgenic brackish medaka is restricted to the reproducing females which synthesize high concentration of estrogen, and no EGFP expression is observed in the liver of embryos, larvae and male fish (FIG. 3). However, when the transgenic brackish medaka fish is exposed to estrogen or estrogen-like chemical substances, EGFP expression can also be induced in the liver of embryos, larvae and male fish (FIG. 3). The expression profile of the introduced EGFP is consistent with the internal expression profile of omChgH in this brackish medaka (X. Chen et al. Ectotoxicol. Environ. Saf. 71:200-208 (2008)), except that omChgH protein is a secretory protein and will transport to ovary through blood circulation after its synthesis in the liver, while EGFP is not a secretory protein and will stay in the liver after its synthesis. As the transgenic brackish medaka female fish express EGFP without any outer induction, it is not easy to differentiate inner EGFP expression and induce EGFP by estrogen-like chemical substances. The non-EGFP-expressing embryo, larvae and male transgenic brackish medaka fish, however, serves as great candidate to monitor estrogen-like activity, which is presented as green fluorescence, especially newly hatched larvae, which swims, does not feed and is the most translucent, is the best stage for estrogen-like activity monitoring.

#### Example 2

##### Estrogen Exposure Experiments

A generalized exposure regimen is established in the present art. FIG. 4 presents an example of using transgenic fish larvae to analyze the estrogen-like activity of a test solution. Exposure of transgenic brackish medaka fish (preferably yolk-sac larvae fish) to test solution (containing estrogen or estrogenic substances) is performed at 28° C. The exposure usually takes 24-48 hours. After exposure, the fish will be deposited on the inner side of a petri-dish cover with a small drop of test solution left. The induced EGFP expression in the fish liver will be observed under fluorescence microscope. Yolk-sac larvae are small, exposure can be carried out in utensils like 96-well plate, 24-well plate, petri-dish and so on to save test reagents. Additionally, newly hatched larvae are the most transparent stage, and the transparency can keep up to 15 days after hatch. After 15 days, the larvae will be gradually less transparent due to the development of muscle. The transgenic fish larvae of 1 to 15 days after hatch have similar estrogen sensitivity (FIG. 5).

In response to different concentrations of estrogen or estrogen-like chemical substances, transgenic fish expresses vary-

ing quantity of EGFP protein which is presented as varying GFP signal intensity under fluorescence microscope. GFP signal intensity increases with estrogen or estrogen-like chemical substances as long as these substances do not reach the concentration to cause lethal toxicity. To quantify the estrogen-like activity of test solution, fluorescent image of fish liver is recorded using a camera connected with the fluorescence microscope. Liver GFP signal intensity can then be measured using some image analysis software such as Metamorph (Universal Imaging). To reduce the interference of fish autofluorescence, only fish liver area was selected for the analysis. The average signal intensity of the manually marked area calculated using the same software was regarded as GFP signal intensity. Fish liver photos recorded using the same image parameter setting can be compared directly using GFP signal intensity, those photos recorded using varying image parameter settings can also be compared as long as each group contain a common reference control.

Considering environmental water samples and other test reagents may have varying salinity and pH values, the effects of different salinity and pH on the sensitivity of transgenic brackish medaka fish to estrogen was investigated. Water of different pH values (e.g., 6.0, 7.0, 8.0 and 9.0) or different salinities (e.g., <5 ppt, fresh water; 15 ppt, brackish water; 30 ppt, common marine water; and 35 ppt, high salinity marine water) was used as test solution, into which the same concentration of estrogen (e.g., 5.1 nM) was added. Newly hatched transgenic brackish medaka fish are exposed to these solutions for a period of time (e.g., 24 hours). Fish livers are recorded and their GFP signal intensity were measured as previously described. Statistical analysis of estrogen induced liver GFP signal intensity proved that neither different pH (ranging from 6.0 to 9.0; FIG. 6) nor different salinity (ranging from 0 to 35 ppt; FIG. 7) affected the sensitivity of this transgenic brackish medaka fish to monitor estrogen-like activity.

### Example 3

#### Estrogen-Like Activity Analysis

##### Natural Estrogens

Estrone (E1), 17 $\beta$ -estradiol (E2) and estriol (E3) are natural steroid estrogens. All the three estrogens can induce liver GFP expression in transgenic brackish medaka larvae of present invention at nM level. Of them, E2 has been intensively studied, and is commonly used as a standard estrogen to evaluate the estrogen-like activity of testing reagents. Using transgenic larvae of present invention, induced GFP signal by E2 can usually be observed within 24 hours, and the GFP signal increases with the increasing concentration of E2. At higher E2 concentrations (e.g., 102 or 1020 nM), induced GFP signal intensity can even be observed within 4 hours after onset of exposure, and the signal increases with exposure time (FIG. 8), and reaches saturation within 24 hours. At low E2 concentrations, however, it will take longer for the induced GFP signal to accumulate to become detectable. Thus, by exposing transgenic brackish medaka fish larvae to different concentrations of E2 for a period of time (such as 24 hours), imaging the GFP expressing liver, and then measuring the induced liver GFP signal intensity using image analysis software, a standard curve correlating GFP signal intensity and E2 concentration can be established (GFP-E2 standard curve; FIG. 9). Using the same regimen, and also include one known E2 concentration exposure as reference, the estrogen-like activity of test reagents can then be evaluated as estrogen

potency equivalent by referring the induced liver GFP signal to that of GFP-E2 standard curve.

##### Synthetic Estrogens

17 $\alpha$ -ethinylestradiol (EE2) and ethinylestradiol 3-methyl ether, the active compounds of contraceptive pills, are synthetic estrogens. EE2 has been widely studied and the estrogenic activity of which is even stronger than that of E2 (K. Kurauchi et al., Environ. Sci. Technol. 39(8):2762-8 (2005); J. Legler et al., Environ. Technol. 36: 4410-4415 (2002); see FIG. 9 and FIG. 10). By exposing the transgenic larvae fish to the same concentration (e.g. in nM) of EE2 and ethinylestradiol 3-methyl ether for the same period of time (e.g. 24 hours), the induced liver GFP signal is almost the same between EE2 and ethinylestradiol 3-methyl ether, which are stronger than that induced by E2. FIG. 10 depicts the dose-dependent response of the liver GFP signal of present invention to EE2.

##### Soymilk Phytoestrogens

Soymilk contains high concentration of phytoestrogens such as genistein, genistin, daidzein and daidzin. Daily consume soymilk is claimed to be good for the health of humans, especially peri- and postmenopausal women, in many aspects (E Lydeking-Olsen et al. Eur J. Nutr. 43(4): 246-257 (2004); M S Kurzer. J. Nutr. 133: 1983S-1986S (2003)). As phytoestrogens like genistein, genistin, daidzein and daidzin are weak estrogens, previous transgenic fish study tried but failed to detect the estrogen activity of genistein (S Scholz et al. Environ Toxicol Chem. 24(10): 2553-2561 (2005)). One recent study found that the estrogen-like activity of pure genistein or genistein in combination with daidzein was not as strong as soymilk (G Rando et al. Toxicol Appl Pharmacol. 237(3): 288-297 (2009)). As pharmaceutical effects of soymilk depends on its estrogen-like activity, it is important to measure the estrogen-like activity of soymilk. Using transgenic medaka larvae fish of present invention, weak GFP signal can be observed in the liver after exposing the larvae to these four phytoestrogens separately at high concentrations for a period of time (e.g. 48 hours). To test the estrogen-like activity of soymilk, phytoestrogens in the soymilk will be firstly extracted out according to procedures we established. Briefly, soymilk is freeze dried and its phytoestrogens are extracted out using methanol. Extract solution is then water evaporated to dryness and phytoestrogens are redissolved in methanol. The concentration of each phytoestrogen can then be quantified by HPLC (high-performance chromatography), and the estrogen-like activity of extracted phytoestrogens can be easily evaluated by measuring induced GFP signal intensity in transgenic fish larvae after exposure to their dilution in water. We found different brand of soymilk has varying estrogen-like activity, and the estrogen-like activities of most soymilks are higher than that of any of the four phytoestrogens alone (e.g., FIG. 11).

##### Industrial Estrogenic Endocrine Disruptors

4-nonyphenol (NP) and bisphenol A (BPA) are well-known endocrine disruptors originated during industry activities. They can mimic estrogen to induce the expression of estrogen-responsive genes such as estrogen receptor, vitellogenin and choriogenin genes (C. Lee et al., J Health Sci, 48(5): 441-445 (2002)). Our previous study also showed that these two chemicals can induce the expression of choriogenin genes in the host organism of present invention (*Oryzias melastigma*) (X. Chen et al. Ectotoxicol. Environ. Saf. 71:200-

208 (2008)). The detection of NP (K. Kurauchi et al., Environ Sci Technol, 39: 2762-2768 (2005)) and BPA (but not NP, Z. Zeng et al., Environ Sci Technol, 39: 9001-9008 (2005)) using transgenic fish have been reported. These two chemicals can also induce GFP expression in the transgenic fish of the present invention.

#### Cosmetic Estrogenic Chemicals

Benzophenone-3 (BP3), 3-(4'-methylbenzylidene-camphor) (4-MBC), homosalate (HMS), octyl dimethyl PABA (OD-PABA), 4-tert-Butyl-4'-methoxydibenzoylmethane (BMDM) and octyl-methoxycinnamate (OMC) are the most commonly used UV blockers in sun creams and cosmetics including lipsticks, skin lotions, hair dyes, shampoos and numerous other products like moldings, jet ink, tires and fabric. Generally, high percentages of UV filters (e.g. up to 10% for BP-3 and 6% for 4-MBC, NR Janjua et al., J Eur Acad Dermatol Venereol. 22(4):456-61 (2008)) are added in the products and a growing use of combinations of different UV filters for absorbing UVA, UVB and UVC light. UV filters enter human blood soon after application (NR Janjua et al., J Eur Acad Dermatol Venereol. 22(4):456-61 (2008)), and present in breast milk of women who used products containing cosmetic UV filters (M Schlumpf et al., Int J Androl. 31(2): 144-151 (2008)). Recent studies showed that many UV filters exhibit estrogenic activity by inducing estrogen receptors (P Y Kunz & K Fent. Toxicol Appl Pharmacol. 217(1): 86-99 (2006); P Y Kunz & K Fent. Toxicol Appl Pharmacol. 234(1): 77-88 (2009)). However, study using transgenic zebrafish failed to detect the estrogen-like activity of UV filters (R Schreurs et al. Arch Toxicol. 76: 257-261 (2002)). Using transgenic *O. melastigma* larvae of present invention, we found BP-3 and 4-MBC can induce GFP expression in the liver after some period of exposure (e.g. 24 hours) at concentration higher than 0.65  $\mu$ M, and the estrogenic activity of BP-3 and 4-MBC mixture is significantly higher than that of these two compounds separately (FIG. 12). Though compounds BMDM, OMC, OD-PABA and HMS, cannot induce the GFP expression in the liver of transgenic *O. melastigma* larvae, the estrogen agonism of BMDM, OMC and OD-PABA can be easily identified by binary exposure with E2 (FIG. 13), and HMS can enhance the estrogenic activity of 17 $\beta$ -estradiol in a dose-dependent manner (FIG. 14).

#### Example 4

##### Anti-Estrogen Activity Analysis

##### Polycyclic Musk

6-acetyl-1,1,2,4,4,7-hexamethyltetraline (AHTN) is one of the most widely used polycyclic musk. This compound is used as a fragrance ingredient in cosmetics and household cleaning products. According to HERA Risk Assessment reports in 2004, AHTN is produced in an annual volume of 1000 to 5000 tons in one Netherland and UK plant, respectively. According to HERA Risk Assessments, AHTN is not toxic to humans basing on present daily consumption. However, the ubiquitous distribution, high accumulation in environment (A Peck et al. Environ Sci Technol. 40: 5629-5635 (2006)), and bioaccumulation in organisms including human beings (H Nakata et al. Environ Sci Technol. 41: 2216-2222 (2007); S Lignell et al. Environ Sci Technol. 42: 6743-6748 (2008)) raise the concern of scientists. AHTN is found to be estrogen antagonist (RHMM Schreurs et al. Environ Sci Technol. 38: 997-1002 (2004); RHMM Schreurs et al. Toxi-

col Sci. 83: 264-272 (2005)). The estrogen antagonist activity of AHTN can also be very easily evaluated using transgenic medaka fish of present invention. Just expose transgenic medaka fish larvae to E2 (e.g. 0.04  $\mu$ M) or E2 in combination with AHTN (e.g. 5  $\mu$ M) for a period of time (e.g. 24 hours), and then measure the GFP signal intensity induced by E2 with/without the presence of AHTN. The weaker GFP signal induced by E2 in the presence of AHTN indicates this chemical is an estrogen antagonist (FIG. 15).

#### Example 5

##### Application of Transgenic *Oryzias Melastigma* Fish to Study Liver Regeneration

Liver plays a central role in metabolic homeostasis by metabolism, synthesis, storage and redistribution of nutrients, carbohydrate, fats and vitamins. At the same time, liver is also the main detoxifying organ of the body through removing waste and xenobiotics by metabolic conversion and biliary excretion. The liver of higher vertebrate, including humans, can precisely regulate its growth and mass and possess high capacity to regenerate after injury. A recent study revealed that fish liver also regenerates after partial hepatectomy in a manner highly similar to that described for mammals (NG Kan et al. FASEB J. fj.09-131730v1 (2009)). As fish share most developmental pathways, physiological and mechanisms and organ systems with humans (AR Cossins and DL Crawford. Nature Rev Genet. 6: 324-333 (2005)), using fish as a model organism for liver regeneration studies may help to understand the molecular and cellular mechanisms underlying liver regeneration. Using transgenic *O. melastigma* fish of present invention, which contains GFP marker for hepatocyte in response to estrogenic endocrine disruptors, will be very useful to reveal the effects of different estrogenic endocrine disruptors on liver regeneration.

Injuries caused by partial hepatectomy of a whole lobe, partial lobe and small scratch surgeries are performed by using protocols employed in previous zebrafish liver regeneration study (N G Kan et al. FASEB J. fj.09-131730v1 (2009)). To study liver regeneration, transgenic adult fish of present invention will be used. Briefly, adult medaka fish are starved for one day and anesthetized in 0.015% tricaine solution before surgery. Fish ventral body is then opened by a 3-4 mm incision and ventral liver lobe is carefully pulled out of the peritoneal cavity and resected at the very base of the lobe with special care. For partial lobe resection and surface scratch surgeries, the ventral lobe is exposed and a small piece of liver tip tissue or a small scratch is introduced using sharp forceps. The remaining liver is then placed back carefully into the peritoneal cavity and the body wall is closed with GLU-ture (Abbott Laboratories). The fish is then placed into fresh water for recovery. Sham-treated fish are also subjected to the same procedure excluding liver resection.

To analyze the effects of endocrine disruptors on liver regeneration process, fish will be treated with/without E2 or other endocrine disruptor(s). Parameters such as whole fish and liver weight are recorded, and liver shape is also recorded depending on its GFP signal using confocal microscope, and its volume can be measured using image analysis software (e.g. Metamorph, Universal Imaging).

Liver morphology changes during liver regeneration is also investigated. Fish liver at different stages of regeneration are dissected out and fixed in 4% paraformaldehyde (PFA)/phosphate-buffered saline (PBS) at 4° C. overnight, washed with PBS containing 0.1% Tween 20, pre-soaked in O.C.T. medium (Tissue Tek) for two hours and then embedded in

O.C.T. and frozen at  $-80^{\circ}\text{C}$ . for more than 3 h before sectioned at  $10\text{ }\mu\text{m}$  thickness using Leica cryostat. Immunohistochemical staining of sections are performed according to standard protocols. Antibodies such as anti-proliferating cell nuclear antigen (PCNA), anti-prox1, anti- $\beta$ -catenin and anti-tubulin are included.

Metabolomic and proteomic alteration of liver hepatocyte are also investigated to understand liver regeneration and the effects of endocrine disruptors on liver regeneration mechanisms. Fish recovering from partial hepatectomy are deeply anesthetized in tricaine solution and livers are dissected and incubated in 0.25% trypsin/PBS solution for 5 min. The liver is triturated to single-cell suspension by gentle pipetting and collected by centrifugation (1000 g, 7 min). Cell suspension is processed to flow cytometry and GFP-labeled hepatocytes are collected and protein extraction is performed according to standard protein extraction protocol. Extracted protein is then separated using NuPAGE Novex Bis-Tris Gels (Invitrogen) at 200V for 15 min, and stained with Colloidal Blue Kit (Invitrogen). Protein bands are cut out and destained with destaining solution (60% water, 30% methanol and 10% acetic acid). After dried in a Speed Vac (Savant), the gel is digested with Trypsin Profile IGD Kit (Sigma). The trypsinized protein digests are dried by Speed Vac and resuspend in 0.1% Formic Acid in water. Peptides are analyzed by HPLC-MS/MS (HPLC: high-performance liquid chromatography, DIONEX Ultimate 300 system; MS/MS: tandem mass spectrometry, BRUKER MicrOTOF-QII). Additionally, iTRAQ (Isobaric tag for relative and absolute quantitation) technique is employed for protein quantitative analysis. The iTRAQ analysis is carried out according to published Nature Protocol (RY Tweedie-Cullen and M Livingstone-Zatchej. Nat. Proc. DIO: 10.1038/nprot.2008.89 (2008)). As a supportative technique, 2-dimension Fluorescence Difference Gel Electrophoresis (2-D DIGE) is also used to analyze the proteomic changes of hepatocytes during liver regeneration. This analysis is performed according to a Nature Protocol (N S Tannu and S E Hemby. Nat Protoc. 1(4): 1732-1742 (2006)).

From the foregoing, although specific embodiments have been depicted and described in detail herein, it will be apparent to those skilled in the relevant art that various modifications, additions, substitutions, and the like can be made without departing from the spirit of the invention and these are therefore considered to be within the scope of the invention as defined in the claims which follow.

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Citation of a reference herein shall not be construed as an admission that such reference is prior art to the present invention. All references cited herein are hereby incorporated by reference in their entirety. Below is a listing of references cited herein:

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## SEQUENCE LISTING

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agagcattcg ccaacatata catttgatgt ggacatcatc ctaataacct catataaaag 4440
gttttttttg accaaaatgt gtacaatatg aggtttttgt gcagattcag ggaaaagatc 4500
actttgtttg tcattgccta ctatcaaaac aaacatttga ggacagataa gttcgagact 4560
gcaaagaccg tgaagggtc tccatgacct ggatgtcaca aaagcctttc attcattcca 4620
acgcaacgac ctgatctggc atttcacgca aaggacagaa tagtcccat gaattacata 4680
aaattgactt aacaaaacac accctgaagc gtttgtgatt gggagctact tggtttgaga 4740
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&lt;210&gt; SEQ ID NO 3

&lt;211&gt; LENGTH: 761

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Oryzias melastigma

&lt;400&gt; SEQUENCE: 3

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tcatttactc gagacctatt agttaaaaa atgtggagga atgagccttt tctttactac 180
gtcgaatatat aaaatttctg agacataaat caattataaa tatacagtat atggcctgat 240
taaaaaaag aaaaaaactt ttcaactgat ttgtaatgc agaaaatcat gcttagcaca 300
accagagcat tcgccaacat atacatttga tgtggacatc atcctaataa cctcatataa 360
aagggttttt ttgacaaaaa tgtgtacaat atgaggtttt gttgcagatt cagggaaaag 420
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actgcaaaga ccctgaaaag gtctccatga cctggatgtc acaaaagcct ttcatctatt 540
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acaaaagcct ttcattcatt ccaacgcaac gacctgatct ggcatttcac gcaaaggaca	180
gaatagtcca catgaattac ataaaattga cttaacaaaa cacaccctga agcgtttgtg	240
attgggagct acttggtttg agagggtggag ttggaagagc atcaaaggta aagacacata	300
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<210> SEQ ID NO 5  
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 <212> TYPE: DNA  
 <213> ORGANISM: Oryzias melastigma

<400> SEQUENCE: 5

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tattgcaaaa acaaaacaag ttgatacata aaggtagcaa catttcttca catttcatgt	180
aaaaaaaaaa aaaaagtctt ttcacatcag tatagcaggt gtgtagatac agttgacaaa	240
atacaacact tccaccttaa tattctatat ggatcaaatg tgactgtttt cagtgaacg	300
tcacgacaat aaagtcacat aatacatttc acttttacac aaatttttac tgtctgtttc	360
tgttcttaaa catacaagca ctgaaaacag agatgaatcc agtataacca aacaactcaa	420
acgacaataa aaaaaacaaa aaaattgttt tattatttta aaatgtttaa aaaaagtcca	480
atttttaaat caaagtaggt caaccatttt taatactgga tcaacaaaca aaaacaatta	540
acaaaaaaaa tcagagttaa tggaaggtaa acacacacat ccgtgaagac aaaaacacaa	600
gattattatt taaaaactga actagacagc ttacttctca gaaatctgag actgtaagga	660
aaactgtttt cctgtgtgct ttcaatttgt aaaattgaaa gatgtcaata aataatttac	720
cctcttgcat ttgaaaaca attgtacttt ctggaagaa tatttggcat aaatgcatgt	780
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<210> SEQ ID NO 6  
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<210> SEQ ID NO 7  
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 <213> ORGANISM: Artificial  
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<400> SEQUENCE: 7

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19

<210> SEQ ID NO 8

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<212> TYPE: DNA

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28

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<212> TYPE: DNA

<213> ORGANISM: Artificial

<220> FEATURE:

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<212> TYPE: DNA

<213> ORGANISM: Artificial

<220> FEATURE:

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<212> TYPE: DNA

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<400> SEQUENCE: 13

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24

<210> SEQ ID NO 14

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<212> TYPE: DNA  
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<212> TYPE: DNA  
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29

<210> SEQ ID NO 21  
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&lt;400&gt; SEQUENCE: 21

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20

<210> SEQ ID NO 22  
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&lt;400&gt; SEQUENCE: 22

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25

<210> SEQ ID NO 23  
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<212> TYPE: DNA  
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<220> FEATURE:  
<223> OTHER INFORMATION: Primer

&lt;400&gt; SEQUENCE: 23

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24

<210> SEQ ID NO 24  
<211> LENGTH: 29  
<212> TYPE: DNA  
<213> ORGANISM: Artificial  
<220> FEATURE:  
<223> OTHER INFORMATION: Primer

&lt;400&gt; SEQUENCE: 24

tgcagcggcc gcctgtgaac cgacagaag

29

<210> SEQ ID NO 25  
<211> LENGTH: 24  
<212> TYPE: DNA  
<213> ORGANISM: Artificial  
<220> FEATURE:  
<223> OTHER INFORMATION: Primer

&lt;400&gt; SEQUENCE: 25

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24

<210> SEQ ID NO 26  
<211> LENGTH: 21  
<212> TYPE: DNA  
<213> ORGANISM: Artificial  
<220> FEATURE:  
<223> OTHER INFORMATION: Primer

&lt;400&gt; SEQUENCE: 26

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21

<210> SEQ ID NO 27  
<211> LENGTH: 23  
<212> TYPE: DNA  
<213> ORGANISM: Artificial

&lt;220&gt; FEATURE:

&lt;223&gt; OTHER INFORMATION: Primer

&lt;400&gt; SEQUENCE: 27

ttacttgtac agctcgctcca tgc

23

What is claimed is:

1. An expression cassette comprising:
  - a reporter nucleotide sequence encoding a reporter protein;
  - an estrogen-responsive 5'-regulatory nucleotide sequence selected from the group consisting of SEQ ID NO:2, SEQ ID NO:3, and SEQ ID NO:4, wherein the regulatory nucleotide sequence is operably linked 5' to the reporter nucleotide sequence to induce expression of the reporter protein in the presence of estrogen or an estrogen-like compound; and
  - a 3'-regulatory region operably linked 3' to the reporter nucleotide sequence.
2. The expression cassette according to claim 1, wherein the reporter protein is an autofluorescent protein.
3. The expression cassette according to claim 2, wherein the autofluorescent protein is selected from the group consisting of an enhanced green fluorescent protein (EGFP), a green fluorescent protein (GFP), a blue fluorescent protein (BFP), a yellow fluorescent protein (YFP), and a red fluorescent protein (RFP).
4. The expression cassette according to claim 1, wherein the 3'-regulatory region comprises an mRNA polyadenylation signal.
5. The expression cassette according to claim 4, wherein the mRNA polyadenylation signal is the 3'-flanking region of a choriogenin H gene of a medaka fish selected from the group consisting of *Oryzias melastigma* and *Oryzias latipes*.
6. The expression cassette according to claim 5, wherein the 3'-flanking region of the choriogenin H gene comprises the 3'-UTR (untranslated region) and the polyadenylation tail sequences of the choriogenin H gene.
7. The expression cassette according to claim 1, wherein the 3'-regulatory region comprises the nucleotide sequence of SEQ ID NO:5.
8. A host cell transduced with the expression cassette according to claim 1.
9. A transgenic cell of an aquatic animal, said transgenic cell comprising at least one expression cassette according to claim 1.
10. The transgenic cell according to claim 9, wherein the aquatic animal is selected from the group consisting of an *Oryzias* species and a *Danio* species.
11. The transgenic cell according to claim 10, wherein the *Oryzias* species is selected from the group consisting of *Oryzias melastigma* and *Oryzias latipes*.
12. An expression cassette comprising:
  - a reporter nucleotide sequence encoding a reporter protein;
  - an estrogen-responsive 5'-regulatory nucleotide sequence isolated from a medaka fish, wherein the regulatory nucleotide sequence is operably linked 5' to the reporter nucleotide sequence to induce expression of the reporter protein in the presence of estrogen or an estrogen-like compound; and
  - a 3'-regulatory region operably linked 3' to the reporter nucleotide sequence, wherein the 3'-regulatory region comprises the nucleotide sequence of SEQ ID NO:5.
13. The expression cassette according to claim 12, wherein the 5'-regulatory nucleotide sequence is isolated from a brackish medaka fish of the species *Oryzias melastigma*.
14. The expression cassette according to claim 13, wherein the 5'-regulatory nucleotide sequence is derived from the promoter region of a choriogenin H gene of *Oryzias melastigma*, said choriogenin H gene having a nucleotide sequence of SEQ ID NO:1.
15. The expression cassette according to claim 12, wherein the reporter protein is an autofluorescent protein.
16. The expression cassette according to claim 15, wherein the autofluorescent protein is selected from the group consisting of an enhanced green fluorescent protein (EGFP), a green fluorescent protein (GFP), a blue fluorescent protein (BFP), a yellow fluorescent protein (YFP), and a red fluorescent protein (RFP).
17. A host cell transduced with the expression cassette according to claim 12.
18. A transgenic cell of an aquatic animal, said transgenic cell comprising at least one expression cassette according to claim 12.
19. The transgenic cell according to claim 18, wherein the aquatic animal is selected from the group consisting of an *Oryzias* species and a *Danio* species.
20. The transgenic cell according to claim 19, wherein the *Oryzias* species is selected from the group consisting of *Oryzias melastigma* and *Oryzias latipes*.

\* \* \* \* \*